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Chitin degradation in C. japonicus

Systems analysis of the Glycoside Hydrolase family 18 enzymes from *Cellvibrio japonicus* characterizes essential chitin degradation functions

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ABSTRACT

Understanding the strategies used by bacteria to degrade polysaccharides constitutes invaluable tool biotechnological for applications. Bacteria are major mediators of polysaccharide degradation in however the complex mechanisms used to detect, degrade, and consume these substrates are not well understood, especially for recalcitrant polysaccharides such as chitin. It has been previously shown that the model bacterial saprophyte Cellvibrio japonicus is able to catabolize chitin, but little is known about the enzymatic machinery underlying this capability. Previous analyses of the C. japonicus genome and proteome indicated the presence of four family 18 Glycoside Hydrolase enzymes, and studies of the proteome indicated that all are involved in chitin utilization. Using a combination of in vitro and in vivo approaches, we have studied the roles of these four chitinases in chitin bioconversion. Genetic analyses showed that only the chil8D gene product is essential for the degradation of chitin substrates. Biochemical characterization of the four enzymes showed functional differences and synergistic effects during chitin degradation, indicating non-redundant roles in the cell. Transcriptomic studies revealed complex regulation of the chitin degradation machinery of C. japonicus and confirmed the importance of CjChi18D and CjLPMO10A, a previously characterized chitin-active enzyme. With this systems biology approach, we deciphered the physiological relevance of the GH18 enzymes for chitin degradation in *C. japonicus*, and the combination of *in vitro* and *in vivo* approaches provided a comprehensive understanding of the initial stages of chitin degradation by this bacterium.

Chitin, a linear polymer of $\beta(1-4)$ -linked N-acetylglucosamine, is the second most abundant polysaccharide on earth after cellulose and a major source of fixed carbon and nitrogen, especially in the oceans (1). The rate of chitin degradation has been calculated to match the rate of synthesis resulting in little net accumulation of chitin in the environment, suggesting that the microorganisms that derive nutrition from this polysaccharide are able to degrade it with high efficiency (2). Moreover, chitin is increasingly being recognized as an important feedstock for the production of renewable chemicals, and chito-oligosaccharides are being investigated for several biomedical applications (3-6). The strategies used by microbes for efficient chitin bioconversion are gaining interest, as current chemical methods used for the industrial processing of chitin are inefficient and wasteful (7,8). Therefore, the

combination of environmental importance and industrial/medical relevance has renewed interest in the bioconversion of chitin (9,10).

Depolymerization of chitin-rich substrates has been studied predominately from biochemical and structural perspectives, and as a consequence there is considerable understanding on certain mechanistic aspects of enzymatic degradation (11). Chitin deconstruction shares many features with cellulose degradation, which is not surprising given that chitin has a crystalline structure similar to that of cellulose and differs chemically via an N-acetyl substitution at the C2 carbon (12). Specifically, chitin depolymerization is achieved through a concerted effort of endo- and exo-acting enzymes to reduce polymer length to short oligosaccharides that are then converted to Nacetylglucosamine monomers hexosaminidases for entry into cellular metabolism (Fig. 1) (6). Additionally, the action polysaccharide monooxygenases of (LPMO) in aerobic microbes promotes chitin degradation by acting on crystalline regions of the substrate (13-15).

Physiological and genetic aspects of chitin degradation, especially for terrestrial bacteria, are not well characterized. One reason for this lack of knowledge is the multiplicity of chitinolytic enzymes in many, but not all, chitindegrading bacteria, which makes functional analysis of the individual enzymes challenging. While several model organisms have been used to study chitin degradation (16-18), one that is emerging as a powerful model, due to the available systems biology tools, is the saprophytic Gram-negative bacterium Cellvibrio japonicus (19,20). This bacterium is a potent chitin degrader (13), and has a suite of nine encoding enzymes with predicted genes functions in chitin degradation Specifically, the C. japonicus genome is predicted to encode four chitinases from the Glycoside Hydrolase family 18 (GH18), one GH19 chitinase, two hexosaminidases (GH20), one chitosanase (GH46), and one chitin-specific LPMO (Auxiliary Activity, AA10) (21). Secretome analysis (22) has shown that the four GH18s, and to a much lesser extent the GH19, are abundant during growth on chitin. Generally, the GH18s and LPMOs are considered key

enzymes for bioconversion of crystalline chitin. The physiological role of GH19 enzymes in chitin degradation is less clear (23,24).

The possession of large numbers of Carbohydrate Active Enzymes (CAZymes) (25) is a hallmark feature of C. japonicus. There is increasing evidence that the CAZymes of C. japonicus that belong to the same GH family are not functionally redundant, but have unique physiological functions (14,26). In the current study, the physiological roles of the four C. japonicus GH18 chitinases were determined focusing on the initial stages of chitin degradation. Sequence analysis indicates that CiChi18B CiChi18C and CiChi18D are secreted enzymes containing at least one chitin-specific carbohydrate-binding module (CBM), whereas CiChi18A is a single domain protein that likely is membrane anchored (Fig. 2). Through RNAseq analysis, we determined that these four chitinases were highly up-regulated while chitin was being used as the sole carbon source. Combinatorial mutational analyses determined that only the chi18D gene product is essential chitin degradation. for Biochemical characterization of the catalytic domains of these four GH18 chitinases indicated that CiChi18D is substantially more active towards insoluble chitin than the other GH18 chitinases. underpinning its importance in chitin degradation. The functional insights into the stages of chitin degradation saprophytic bacteria that are provided in this study have the potential to accelerate the development of industrial chitin bioconversion strategies.

Results

CjChi18D is essential for the degradation of α -chitin

According to previous sequence analysis (21), C. japonicus possesses four GH18 chitinases (CjChi18A, CjChi18B, CjChi18C and CjChi18D). A recent proteomic study (22) detected the four enzymes in the secretome of C. japonicus growing on α - and β -chitin, which suggested the importance of these enzymes for efficient chitin degradation. To elucidate the physiological relevance of these enzymes, we generated a suite of GH18 deletion mutants and assessed their fitness on insoluble chitin

substrates, including an environmentally relevant substrate, crab shell.

Wild type and GH18 deletion mutant strains all grew well in defined media with either glucose (Glc) or *N*-acetylglucosamine (GlcNAc) as the sole source of carbon (Fig. S1). A Δgsp mutant, which lacks the entire 9.4 kb gsp operon encoding the Type II Secretion System that is needed for secretion of most enzymes, was also able to grow on these monosaccharides, as previously demonstrated (13,27). When the GH18 single deletion mutants were grown using insoluble α -chitin or crab shell as the sole carbon source distinct phenotypes emerged (Fig. 3). When α -chitin was the sole source of carbon, the $\Delta chi18D$ mutant was unable to grow. The other three single deletion strains ($\Delta chi18A$, $\Delta chi18B$, and $\Delta chi18C$) all had similar growth rates and maximum optical densities (OD) as the wild type (Table S1A). Interestingly, the $\Delta chi18B$ and $\Delta chi18C$ single mutant strains had more protracted lag phases than the wild type strain when using α-chitin as carbon source (Fig. 3A). When using crab shells as the only source of carbon, the $\Delta chi18D$ mutant was unable to grow while the single mutants of $\Delta chi18A$, $\Delta chi18B$, and $\Delta chi18C$ displayed growth similar to wild type (Fig. 3C and Table S1B).

To test for functional redundancies between the GH18 chitinases, we generated all combinations of double deletion mutants, and then assessed growth using α -chitin or crab shells as the sole carbon source. The double mutant $\Delta chi18B$ $\Delta chi18C$ resulted in a slower growth rate and a longer lag phase when grown on α -chitin in comparison to either of the single deletion mutants or the wild type (Fig. 3B). The growth rate of the $\Delta chi18B$ $\Delta chi18C$ double mutant was reduced 37% compared to wild type (**Table S1A**). The $\Delta chi18A$ $\Delta chi18B$ double mutant grew like the $\Delta chi18B$ single mutant while the $\Delta chi18A$ $\Delta chi18C$ double mutant grew like the $\Delta chi18C$ single mutant. The $\Delta chi18A$ Δchi18B Δchi18C triple mutant recapitulated the growth defect observed in the $\Delta chi18B$ $\Delta chi18C$ double mutant in terms of the growth rate, lag phase, and the maximum OD (Fig. 3B and Table S1A). These results suggest that the chi18B and chi18C gene products have nonredundant functions during degradation of α -chitin, whereas the *chi18A* gene product does not play a rate-limiting role.

Interestingly, double the mutants exhibited wild type-like phenotypes when grown using crab shells as the sole carbon source (Fig. **3D**). The growth rates of the $\Delta chi18B$ and $\Delta chi18C$ single mutants and the $\Delta chi18B$ $\Delta chi18C$ double mutant were similar to the wild type strain when grown using crab shells. For all tested strains the growth rates when using crab shells as the sole carbon source were substantially reduced compared to α -chitin. The results of experiments using β -chitin (Fig. S2) were very similar to the results obtained with α chitin.

CjChi18D is a potent secreted chitinase

Sequence alignment of the catalytic domains of the four chitinase showed large sequence variation. CjChi18A and CjChi18D share higher sequence identity (32%), while CiChi18C and CiChi18D showed the lowest identity (19%), as determined by BLAST alignment (28). As predicted by LipoP (29) and SignalP (30) software tools, all four of the C. japonicus GH18 chitinases have a signal sequence, however CiChi18A has a SPaseIIcleaveable sequence and is predicted to be an outer membrane associated lipoprotein. The CiChi18B, CiChi18C and CiChi18D enzymes have a SPaseI-cleaved sequence (29) and are predicted to be secreted. The domain structure of the four GH18 chitinases is summarized in Fig. 2. The LipoP/SignalP software predictions are in agreement with a proteomics study that showed the occurrence of all four GH18 chitinases in the secretome (22).

To further assess the contribution of individual GH18 enzymes as effectors of chitin degradation, we used our suite of GH18 mutants to assess secreted activity using colloidal chitin plate assays (**Fig. 4** and **Table S2**). A wild type strain generated a robust zone of clearing while a Δgsp secretion-deficient mutant generated no zone of clearing. We observed a similarly striking phenotype with the $\Delta chi18D$ single mutant strain, which also generated no zone of clearing. The remaining GH18 single mutants displayed approximately wild type zones of

clearing. Interestingly, relative to the wild type, the $\Delta chi18B$ $\Delta chi18C$ double mutant showed a clear reduction in the size of the clearing zone. The $\Delta chi18A$ $\Delta chi18B$ $\Delta chi18C$ triple mutant displayed a zone of clearing similar to the $\Delta chi18B$ $\Delta chi18C$ double mutant. The reduced chitin-degrading capacities of the $\Delta chi18B$ $\Delta chi18C$ double mutant and the $\Delta chi18A$ $\Delta chi18B$ $\Delta chi18C$ triple mutant are in agreement with the observed growth defects of these strains when grown on α -chitin.

The GH18 chitinases have different activities towards chitin and $(GlcNAc)_6$.

To further investigate the features of the four GH18 chitinases, they were cloned, expressed, and purified for biochemical characterization. Despite substantial efforts, soluble full-length multi-domain CiChi18B, CiChi18C, and CiChi18D proteins could not be obtained. Therefore comparative biochemical analysis was conducted with over-expressed catalytic domains. All four chitinases were able to degrade α -chitin, although with greatly varying efficiency, but clearly highest for CiChi18D (Fig. 5). As expected (31), versions of CiChi18B and CiChi18D containing their CBM5 domain gave higher yields compared to their respective solitary catalytic domains. The catalytic domain of CiChi18C gave higher yields than the catalytic domains of CiChi18A and CiChi18B, but all were poor in chitin degradation compared to CiChi18D.

Analysis of products generated during degradation of α-chitin showed that CjChi18A_{cat} initially produced both GlcNAc and (GlcNAc)₂. The product profiles for the other variants (CjChi18B_{cat}, CjChi18B_{cat+CBM5}, CjChi18C_{cat}, CjChi18D_{cat}, and CjChi18D_{cat+CBM5}) showed mainly (GlcNAc)2 and smaller amounts of GlcNAc (Fig. S3), as is usual for GH18 chitinases. It is noteworthy that the product profile of CjChi18Bcat was different from CjChi18B_{cat+CBM5} after 48h, where GlcNAc is the dominating product for CiChi18B_{cat}, while (GlcNAc)₂ is the dominating product for CiChi18B_{cat+CBM5} (Fig. S3). For CiChi18A_{cat}, only GlcNAc was detected after 48 h, indicating an N-acetylhexosaminidase activity for this enzyme.

Synergy experiments (Fig. 6) confirmed the dominating role of CiChi18D in chitin bioconversion and revealed synergistic effects for various enzyme combinations, such as when CiChi18A_{cat} with CiChi18B_{cat}, CjChi18B_{cat} with CjChi18C_{cat}, and CjChi18B_{cat} with CiChi18Dcat. These synergistic effects indicate that the various chitinases must have different functionalities. As expected on the basis of the data presented above, the presence of CjChi18A_{cat} shifted the product profile towards a higher GlcNAc/(GlcNAc)₂ ratio. The effect of adding CjLPMO10A was generally small, but seemed slightly larger when combined with the individual catalytic domains of CiChi18A, CiChi18B, and CiChi18C, compared to CjChi18D. While CjChi18D alone was the most potent individual enzyme, the highest chitin solubilization yields were obtained when CiChi18D was paired with other chitinases.

Investigation of enzyme activity towards soluble chito-oligosaccharides can give insight into preferred substrate binding modes and intrinsic catalytic rates. Thus, activity towards the soluble chito-oligosaccharide (GlcNAc)₆ was determined for all four GH18 catalytic domains (Fig. 7A). The results indicated that CiChi18C_{cat} had the highest activity against (GlcNAc)₆ with an initial rate of $300 \pm 8 \text{ min}^{-1}$. CjChi18A_{cat} had an initial rate of $118 \pm 1 \text{ min}^{-1}$, while CiChi18D_{cat} and CiChi18B_{cat} had initial rates of 44.5 ± 2.6 and 27.0 ± 0.8 min⁻¹, respectively. The product profile obtained for the enzymes shortly after initiation of the (GlcNAc)₆ hydrolysis reactions (2 min reaction time) showed striking differences between the chitinases (Fig. 7B). CiChi18Acat yielded all possible product types ((GlcNAc)₁₋₅), where GlcNAc was the dominating product (97.8 ± 7.3) μ M) followed by (GlcNAc)₂ (62.3 ± 2.5 μ M), $(GlcNAc)_3 (47.7 \pm 1.7 \mu M)$ and $(GlcNAc)_4 (33.9)$ \pm 1.5 μ M). Quantification of (GlcNAc)₅ was not possible due to co-elution with (GlcNAc)₆. CiChi18B_{cat} produced mainly (GlcNAc)₂ and (GlcNAc)₃, with minor amounts of (GlcNAc)₄. CjChi18C_{cat} produced (GlcNAc)₂₋₄ CiChi18D_{cat} produced mainly (GlcNAc)₂ and (GlcNAc)₄. Chito-oligosaccharides longer than (GlcNAc)₆ were observed in the CiChi18D_{cat} reaction (Fig. 7B and Fig. S4), indicating that this enzyme has transglycosylating activity.

Reactions using (GlcNAc)₂ as a substrate showed rapid conversion by *Cj*Chi18A_{cat}, whereas only trace amounts of the monomer were detected for the other three chitinase (results not shown).

C. japonicus GH18 chitinase genes are highly expressed during chitin utilization.

To obtain further insight into the regulation of chitinase expression and to complement the previous secretome study (22), we used RNAseq to determine changes in gene expression during growth using α -chitin relative to growth using glucose. Analysis of samples from the exponential growth phase revealed significant up-regulation of 73 CAZyme genes (Table S3A). The seven most strongly upregulated genes were all related to chitin bioconversion, including the LPMO (6.7-fold, log₂ scale), the hex20B hexosaminidase (4.1fold), the nag9A gene (3.5-fold) that encodes a putative *N*-acetylhexosamine 6-phosphate deacetylase, and the four GH18 chitinases in the following order chi18D> chi18C > chi18A > chi18B, the fold change (log₂ scale) being 6.7fold, 5.1-fold, 3.9-fold and 3.6-fold, respectively (Fig. 8 and Table S3A). Several other genes putatively involved in chitin bioconversion were also up-regulated, albeit to a lesser extent, including the GH19 chitinase (2.2-fold, log₂ scale), the other GH20 hexosaminidase (hex20A; 2.7-fold), the GH46 chitosanase and two polysaccharide deacetylases (2.6-fold, 1.6-fold, and 1.4-fold, respectively).

The comparison of gene expression during early stationary phase yielded similar results (Fig. S5A and Table S3B). This analysis revealed the up-regulation of 47 CAZyme genes, of which seven are implicated in chitin degradation (lpmo10A, chi18D, chi18C, chi18B, nag9A, hex20A, and chi19A). While the expression data, in particular for the exponential comparison, showed a strong up-regulation of chitin-relevant genes, there was also upregulation of a wide variety of CAZyme genes associated with the degradation of other polysaccharides such as starch, xylan, cellulose, pectin, arabinanan, mannan, β-glucan and xyloglucan. Therefore, the regulation of the chitin response resembles the general response previously observed for cellulose (14). When

comparing the transcriptomes of the exponential and the stationary phase during growth on chitin, none of the significant changes in expression concerned CAZyme genes (Fig. S5B).

Discussion

A previous report demonstrated that C. japonicus can grow using both purified chitin and unrefined crab shells as a sole nutrient source (13). Additionally, a recent proteomic study found several secreted GH18 chitinases, suggesting a robust response to chitin (22). It has remained unknown, however, how these GH18 chitinases contribute to the degradation of chitin, and to what extent the four GH18 genes in the C. japonicus genome are equivalent. In this study, through an implementation of transcriptomic, genetic, and biochemical approaches, we assessed the physiological functions of the four GH18 enzymes. The synthesis of in vitro, in and in silico data provides a of comprehensive understanding chitin degradation by C. japonicus.

The architecture of the family GH18 chitinases indicates differential biological roles

Analysis using LipoP (29) predicts that the CiChi18A enzyme is a lipoprotein with a signal sequence cleaved by signal peptidase II (SPII). The presence of a glutamine in position +2 after the cleavage site of SPII further suggests that CiChi18A resides attached to the membrane (32-35).Interestingly, CiChi18A was found in high amounts in a recent secretome study (22), however the precise location of this enzyme remains undefined. It is likely that CiChi18A functions in degrading short chito-oligosaccharides, as the absence of CBMs limits the association of this enzyme with polymeric chitin (31,36). Such a function is supported by the biochemical data for CiChi18A showing high activity on soluble substrates and an ability to produce monomers.

Analysis of the CjChi18B, CjChi18C and CjChi18D enzymes indicated a signal peptide that is cleaved by signal peptidase I (29). The growth defect on chitin observed in the Δgsp mutant suggest that these three enzymes are transported to the extracellular space by the Type II Secretion System (37). The CjChi18B, CjChi18C, and CjChi18D enzymes all possess a

CBM5 and a CBM73 domain (38), however the orientations of these domains differ between the enzymes (**Fig. 2**). As found with many CAZymes, the *Cj*Chi18B, *Cj*Chi18C, and *Cj*Chi18D chitinases contain serine rich regions separating the individual domains, which likely act as flexible linkers (39).

Analysis of the domain organization of the C. japonicus chitinases, functional data, homology modelling of their 3-D stuctures (Fig. S6), and comparison with well-characterized GH18 chitinases from Serratia marcescens (18) all indicate that the C. japonicus chitinolytic system represents a suite of complementary activities. Available data from known processive chitinases show that there are hallmarks specific for processive enzymes, involving an $\alpha+\beta$ domain in the structure, and conserved aromatic amino acids lining the catalytic cleft (40-43). Structural modelling of the C. japonicus GH18 chitinases and sequence alignment with known processive chitinases (Fig. S7), show that CjChi18B and CiChi18D have specifications. CiChi18B has a large $\alpha+\beta$ domain (~110 amino acids) and an "tunnel-like" substrate-binding cleft, while CjChi18D has a much smaller $\alpha+\beta$ domain (~60 amino acids) and a more open active site cleft. Compared to the chitinases from S. marcescens (Fig. S6), the structural model of CiChi18B shows a more closed active site than exo-processive SmChiA and SmChiB, while CjChi18D seems to have a more open active site cleft than these two Serratia enzymes. CiChi18A has the $\alpha+\beta$ domain (~65 amino acids), but lacks most of the aromatic residues; the most similar S. marcescens enzyme seems to be SmChiD, which, like CiChi18A, primarily produces monomers (44) CjChi18C does not have features characteristic for processivity and is more similar to SmChiC, which is a non-processive endo-chitinase, displaying an open substrate binding cleft that is characteristic for nonprocessive glycoside hydrolases (36).

The GH18 enzymes of Cellvibrio japonicus have divergent functionalities but are reliant on CjChi18D

The growth and chitinase secretion data strongly pointed to the *chi18D* gene product being essential for the degradation of chitinous

substrates (Fig. 3, Fig. 4, and Fig. S2). These results reinforce two features previously described for C. japonicus, specifically, that secreted enzymes are essential for recalcitrant polysaccharide degradation and that single CAZymes elicit major physiological effects (27,45-47). Other individual enzymes did not seem essential for growth, but the $\Delta chi18B$ $\Delta chi18C$ double deletion mutant did show reduced growth and reduced chitin conversion efficiency (Fig. 3 and Fig. 4), indicating that these two chitinases have complementing functionalities and that they contribute to chitin conversion but are not essential. The absence of growth defects of the multiple mutants while growing using crab shells is likely a consequence of the slow overall growth rate.

Biochemical characterization of the catalytic domains revealed clear functional differences (Fig. 5, Fig. 7, Fig. S4, and Table **S4**), and showed that the GH18 enzymes act synergistically during the degradation of chitin (Fig. 6). Enzyme synergy during the course of chitin degradation has been described previously for S. marcescens, and is generally ascribed to collaboration between endo- and exo-acting enzymes, where endo-acting enzymes generate chain ends that are substrates for exo-acting enzymes that usually show processive action (48). The synergistic effects observed for the best combinations of C. japonicus chitinases (Fig. 6) are of similar size as the effects observed for the S. marcescens enzymes (49,50). It is worth noting that the clearest synergy was observed when combining CiChi18B, for which there are strong indications of exo-processive action, and CiChi18C or CiChi18D, which both could be endo-acting enzymes (vide infra). Previous studies with β-chitin have shown that supernatants of chitin-grown marcescens and chitin-grown C. japonicus have similar chitinolytic power (22).

The GH18 chitinases exhibit processivity during chitin hydrolysis

Although it is difficult to determine the degree of processivity from standard substrate degradation experiments, such experiments do provide indications. During degradation of crystalline chitin, processive enzymes tend to generate high (GlcNAc)₂/GlcNAc ratios (43,51).

Furthermore, the degradation of (GlcNAc)₆ by processive enzymes will yield a high (GlcNAc)₂/GlcNAc)₄ ratio since such enzymes will process the chitohexaose into three chitobiose moieties, rather than releasing the chitotetraose and chitobiose into solution after a single hydrolytic event (as would be expected for a non-processive enzyme). Of the GH18 chitinases of C. japonicus, only CjChi18B showed these high ratios (Table S4), indicating that this enzyme is the most processive, as also suggested by the structural model discussed above (Fig. S6). The other three enzymes did not show clear signs of processivity. CiChi18C displays all hallmarks of a non-processive enzyme, and CiChi18A is unique as it primarily produces monomers, which is not really compatible with processivity since the repeating unit in a chitin chain is a dimer. These latter two enzymes showed the highest specific activities towards the hexamer, suggesting that they primarily act on soluble substrates. On the basis of the experimental data it is difficult to judge whether the four enzymes are endo- or exoacting, and it should be noted that mixed endo-/exo-modes do occur (52). CjChi18B appears to be exo-processive, while CjChi18C looks to be endo-non-processive. The situation CiChi18A and CiChi18D is less clear, since the deep but still relatively open active site clefts of these enzymes are compatible with both endoand exo-action, and since the biochemical data were not conclusive regarding processivity.

The chitinolytic machinery of C. japonicus is highly responsive to the presence of chitin substrates

There was significant up-regulation of the predicted *C. japonicus* genes encoding chitin-active proteins (GH18, GH19, GH20, GH46, and AA10 families) both during exponential and stationary phase (**Table S3**). The up-regulation of a number of other CAZyme genes not involved in chitin degradation was striking, but not surprising. *C. japonicus* has been shown to have both substrate-dependent and growth rate-dependent control of CAZyme gene expression (14). In regards to the substrate-specific response, *C. japonicus* seems to have two variations, one that is general (where diverse CAZyme genes are up-regulated), and

one that is specific (where only substrate-specific CAZymes genes are up-regulated). The response observed for cellulose is an example of the former (14), and the response for xylan is the latter (46). The regulatory network for the chitinolytic response appears to be analogous to the cellulose response, which is not surprising given the similarities of the substrates.

The expression data also indicated that the most up-regulated chitin-relevant gene the chitin-specific encodes LPMO. CiLPMO10A. It has been shown that this enzyme will act synergistically with chitinases (13) but in the present study, the effects of adding the LPMO to the chitinases was limited (Fig. 6). This is likely an effect of the substrate used in this study, which was balled milled into fine particles. It has previously been shown that such ball milling reduces the crystallinity of the substrate, hence reducing the activity of the LPMO (53).

Final remarks

The proposed model (Fig. 1) for crystalline chitin degradation by C. japonicus, based on in vitro, in vivo, and in silico analyses that the four GH18 chitinases and CiLPMO10A synergistically chitin. CiLPMO10A plays an important role as indicated by the transcriptomic data presented here and previously published biochemical data (13). CiChi18D, being the most powerful chitindegrader, likely also acts on the more recalcitrant parts of the substrate, releasing chito-oligosaccharides. CiChi18B and CiChi18C act on the more accessible chitin fibers converting them smaller chitointo oligosaccharides, while CiChi18A converts chito-oligosaccharides to N-acetylglucosamine.

Experimental procedures *Growing conditions*

Cellvibrio japonicus strains were grown using MOPS (3-(N-morpholino)propanesulfonic acid)) minimal media (54) containing 0.2% wt/vol glucose, 0.5% wt/vol N-acetylglucosamine, 0.25% wt/vol α -chitin from shrimp shells (Sigma, Aldrich), 0.2% wt/vol β -chitin from squid pen, or 4% wt/vol crab shells from Callinectes sapidus as the sole source of carbon. The following protocols were developed

to prepare the chitin containing substrates. The coarse flakes of α -chitin were sieved through the top piece of a 130 mm Buchner polypropylene filter to homogenize the particle size (diameter of ca. 4 mm). β-chitin from squid pen was manually ground in a ceramic mortar and pestle and then passed through a plastic filter with 4 mm diameter holes. For the crab shells, any membranous material was discarded and the shells were then thoroughly washed and rinsed. Clean and air-dried shells were manually ground and then sieved through a 4 mm diameter filter. To remove any generated crab shell dust, the filtrate was rinsed through a Buchner polypropylene filter. The shell pieces were subjected to autoclave sterilization. Escherichia coli strains were grown using lysogenic broth (LB) (55). For growth analysis studies, strains were initially grown for 24 h in 5 mL of MOPSglucose medium. Then, an 18 mm test tube with 5 mL MOPS-chitin broth was inoculated with a 1:100 dilution of the prepared overnight cultures. All cultures were incubated at 30 °C with an aeration of 225 RPM. For insoluble substrates growth was measured as a function of the optical density at 600 nm (OD₆₀₀) with a Spec20D+ spectrophotometer Scientific) using biomass containment as needed (56). In growth experiments with glucose or Nacetylglucosamine growth was monitored using a Tecan M200Pro microplate reader at 600 nm (OD₆₀₀) (Tecan Trading AG, Switzerland). All experiments were performed in growth biological triplicates. Plate media were solidified with 1.5% agar. When required, kanamycin was used at a concentration of 50 µg/mL.

Generation of deletion mutants

Deletion mutants of the family GH18 chitinases of *C. japonicus* were made and verified using previously published protocols (27,45). A suicide vector was generated by cloning ±500 bp up- and down-stream from the gene to be deleted into the pK18*mobsacB* vector (57) at the EcoRI and the XbaI sites. The pK18*mobsacB* plasmids to make the deletions of genes *chi18A*, *chi18B* and *chi18C* were synthetized by GeneWiz (South Plainfield, NJ), while the plasmid to make the deletion of *chi18D* was amplified by PCR and assembled via Gibson assembly (58). All vectors were

electroporated into $E.\ coli\ S17\ \lambda_{PIR}$. Through triparental mating the deletion plasmid was conjugated into $C.\ japonicus$ using an $E.\ coli$ helper strain containing the plasmid pRK2013 (59). Recombinant colonies were selected using kanamycin selection, with a counter selection carried out via sucrose toxicity. The deletion mutants were confirmed by PCR, and the primers used for the construction and verification of $C.\ japonicus$ mutants are listed in **Table S5**.

Visualization of colloidal chitin degradation

Colloidal chitin was prepared as described in (60). Colloidal chitin plates were made using MOPS defined media supplemented with 2% (w/v) colloidal chitin and 0.2% (w/v) glucose. To assay for chitin degradation, 10 µL of overnight cultures of the *C. japonicus* strains to be analyzed were spotted onto the chitin plate and incubated for 4 d at 30 °C. Then, plates were stained with a 0.1% (w/v) Congo red solution for 10 min followed by 10 min destaining with a 1M NaCl solution, as described previously for detection of degradation of carboxyl methyl cellulose (27,45,56).

Transcriptomic analysis

analysis transcriptomic conducted for C. japonicus grown using glucose or α-chitin. To prepare the samples, the protocol described in Gardner et al. (14) was followed. Briefly, 35 mL of cell culture were aliquoted into 50 mL conical tubes with 5 mL of a stopping solution made with ethanol and saturated phenol (19:1). The cells were pelleted by centrifugation at 8000 g for 5 min at 4 °C. The supernatant was removed and the cell pellets immersed in a dry ice/ethanol bath for 5 min. The frozen cell pellets were then stored at -80 °C. For each carbon source, samples were taken in biological triplicate and at two time points: the beginning of the exponential phase $(0.1>OD_{600}>0.2)$ and the beginning stationary phase. GeneWiz performed RNAseq using an Illumina HiSeq2500 (50bp single-reads; >10 million reads per sample), and sequence data was analyzed using CLC Genomics Workbench (QIAGEN). The log₂ transformation and a Student's T-test were used for the comparative analysis of C. japonicus gene expression. An

adjusted p-value > 0.01 and a log₂ fold change > 2 were selected as significance cut-off parameters. RNAseq data for glucose grown *C. japonicus* cells (GSE90955) was used as done previously (26). The chitin-specific RNAseq data that was generated for this study can be found in the NCBI GEO database (GSE108935).

Bioinformatics analysis

We determined the predicted CAZy domains presented using the <u>Database</u> for Automated <u>Carbohydrate-active</u> enzyme <u>ANn</u>otation (DbCAN)(38). Using LipoP (29) and SignalP (30), we determined the putative location of the chitinolytic enzymes. Three-dimensional models of the GH18 chitinases were generated using PyMod 2.0 (61).

Cloning and expression of chitinases

Synthetic genes of the four full-length chitinases, optimized for expression in E. coli, were purchased from GeneScript. These genes encoded the catalytic domain for CiChi18A and the full-length protein (without signal peptide) for CiChi18B, CiChi18C and CiChi18D. Primers for amplification of the genes were designed so that a 6xHis-tag followed by a TEV protease cleavage site was introduced at the Nterminus of all proteins and a stop codon was introduced before the C-terminal His-tag encoded in the vector. Using different primers for amplification, genes encoding different versions (Table S6) of the modular proteins were generated, which were cloned into the pNIC-CH vector (62) using ligation independent cloning (63). The DNA sequences of the genes were confirmed by Sanger sequencing. Plasmids were transformed into E. coli BL21 cells. Protein expression was tested by inoculating 50 mL LB + 50 μg/mL kanamycin with 500 μL of an overnight culture. The cultures were grown at 37 °C and when the OD reached 0.6, the cells were induced by adding IPTG to a final concentration of 0.2 mM. After growth overnight at 30 °C, the cells was harvested by centrifugation (6164 g, 12 min, 4 °C) and resuspended in 5 mL 20 mM Tris-HCl, 150 mM NaCl, 5 mM imidazole, pH 8.0. DNAseI and PMSF were added to final concentrations of 1.4 μg/mL and 0.1 mM, respectively, before the cells were lysed by sonication (pulse 3 s on, 3 s off for 3-5 min), using a VC750 VibraCell sonicator (Sonics & Materials, Inc, CT, USA). Following cell disruption, the samples were centrifuged (11814 g, 12 min, 4 °C), after which the supernatant was collected and filtrated (0.22 μ m). Analysis by SDS-PAGE showed that only CjChi18A was soluble. Except for the full-length enzymes, which were not produced in detectable levels, all other enzyme variants were produced in large amounts, but were insoluble.

Production of *CjChi18A* was scaled up to a 500 mL culture following the same protocol as above for production and harvesting. The filtrated supernatant containing *CjChi18A* was used for protein purification by nickel affinity chromatography using a HisTrap HP 5 mL column (GE Healthcare Life Sciences, Uppsala, Sweden) connected to an Äkta pure system (GE Healthcare Life Sciences, Uppsala, Sweden). A continuous imidazole gradient ending at 300 mM imidazole was used to elute bound protein, using a flow rate of 1 mL/min.

CjChi18B_{cat}, CiChi18B_{cat+CBM5}, CjChi18Ccat, CjChi18Dcat, and CjChi18Dcat+CBM5 were produced using a denaturing and refolding method, starting with 500 mL cultures as described above. After harvesting the cells (7025 g, 12 min, 4 °C), the pellet was re-suspended in 20 mL 50 mM Tris, pH 8.0, 0.2 M NaCl and 50% of the cell suspension was used in the further steps. After another centrifugation (18459 g, 10 min, 4 °C) the pellet was resuspended in 10 mL 50 mM Tris, pH 8.0, 0.2M NaCl and, after addition of lysozyme (final concentration 200µg/mL) and DNAseI (final concentration 0.1 mM), the samples were incubated on ice for 30 min. The samples were subsequently sonicated (pulse 3 s on, 3 s off, 2-4 min) and inclusion bodies were harvested by centrifugation (18459 g, 10 min), after which the protein pellet was re-suspended in 25 mL washing buffer (20 mM Tris-HCl, 100 mM NaCl, 2% Triton X-100, pH 8.0). The centrifugation and subsequent re-suspension in washing buffer steps were repeated twice and after the final centrifugation, the pellet was dissolved in 5 mL cold denaturing solution (50 mM Tris-HCl, pH 8.5, 100 mM NaCl, 5 M guanidine HCl, 1 mM EDTA, and 20 mM DTT), vortexed for 1 min, and incubated overnight at 4 °C with slow rotation of the sample tubes. The samples were then centrifuged at 9000 g, for 10 min at 4 °C. The supernatant containing the denatured protein was then collected for refolding.

The sample containing the denatured protein (approximately 5 mL) was added to 250 mL cold refolding buffer (100 mM Tris-HCl, 0.4 M L-arginine, 0.5 mM oxidized glutathione, 5 mM reduced glutathione) at a rate of 1 mL/h under intense stirring at 4 °C. After adding all protein, the solution was stirred overnight at 4 °C, before centrifugation (9000 g, 10 min, 4 °C). The supernatant was collected and dialyzed (10 MWCO Snakeskin, Thermo Fisher Scientific, MA, USA) against 2.5 L 50 mM Tris-HCl, 0.1 M NaCl, pH 8.0 overnight. The dialysis solution was changed once, after ca. 6 h. After collecting the dialyzed sample, the protein was purified with nickel affinity chromatography as described above. All refolded proteins stayed soluble during these procedures and the yields of recovered purified soluble protein typically 1-5 mg/L culture.

The presence and purity of the proteins was confirmed by SDS-PAGE, and relevant fractions were pooled before the protein solution was concentrated to 26.4 mg/mL for CjChi18A and 1 - 3.5 mg/mL for the unfolded/refolded chitinases, with concomitant buffer exchange to 50 mM Tris-HCl, 100 mM NaCl, pH 8.0, using Amicon Ultra-15 centrifugal filters with 10 000 NMWL (Merck Millipore, Cork, Ireland). All resulting protein solutions were stable for weeks at 4 °C. Protein concentrations were determined by measuring the absorbance at 280 nm and using the theoretical extinction coefficient (calculated using the ExPASy ProtParam tool available from

https://web.expasy.org/protparam/).

The catalytic domain of CjLPMO10A was expressed and purified as previously described by Forsberg et al. (13).

Activity assays

Standard reactions contained 15 g/L αchitin (extracted from Pandalus borealis, Seagarden, Karmsund, Norway) or 2 mM chitooligosaccharides (MegaZyme, Bray, Ireland), 0.1 mg/mL BSA and 20 mM BisTris, pH 6.5, unless stated otherwise. For chitin degradation 0.5 µM enzyme was used, while for chito-

oligosaccharide degradation 50 nM enzyme was used, unless stated otherwise. Reaction mixtures were incubated in a thermomixer at 30 °C, 800 RPM and enzyme activity was quenched by adding sulphuric acid to a final concentration of 25 mM. A Rezex RFQ-Fast Acid H+ (8%) ionexclusion column (Phenomenex, CA, USA) installed on a Dionex Ultimate 3000 HPLC with UV-detection (194 nm) was used to analyze and quantify degradation products as described by Mekasha et al. (64). Conversion yields were calculated as percentage of the theoretical maximum after correction of the amount of chitin in the reaction for ash and water content. Reactions with (GlcNAc)₆ as substrate were analyzed using a Rezex ROA-organic Acid H+ (8%) ion exclusion column (Phenomenex, CA, USA) installed on a Dionex Ultimate 3000 HPLC, using a column temperature of 65 °C. An 8 µL sample was injected on the column, and the mono/oligosaccharides were eluted isocratically at 0.6 mL/min with 5 mM sulphuric acid as mobile phase. The chito-oligosaccharides were monitored by measuring the absorbance at 194 nm. Standards with known concentrations of GlcNAc (Sigma, MO, USA) and (GlcNAc)₂₋₆ (MegaZyme, Bray, Ireland) were used to determine the concentrations of (GlcNAc)₆ and the degradation products.

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Conflict of interest

This article does not contain any studies using human participants or animals. In addition, the authors declare that they have no conflict of interest.

Author contributions

- **E.C.M.** Generated *C. japonicus* mutants, performed the growth and physiology experiments, generated and analyzed the RNAseq data, and contributed to writing the manuscript.
- **T.R.T.** Cloned and produced the chitinases, performed the biochemical characterization, analyzed the data, and contributed to writing the manuscript.
- **G.V.K.** Participated in planning and analysis of the biochemical characterization of the chitinases and contributed to writing the manuscript.
- **V.G.H.E** Participated in planning and analysis of the biochemical characterization of the chitinases and contributed to writing the manuscript.
- **J.G.G.** Designed the study, provided overall project direction, and contributed to writing the manuscript. All authors read and approved the final submitted version of the manuscript.

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CjChi18D CjChi18B CjChi18C OM transporter OM CjChi18A IM Transporter

Figure 1. Proposed model for chitin utilization in *C. japonicus. CjC*hi18D (green) and *CjLPMO10A* (yellow) work together to disrupt the crystalline structure of chitin and degrade less accessible chitin fibers, while *CjC*hi18B (magenta) and *CjC*hi18C (brown) are acting on the more accessible chitin fibers to produce chito-oligosaccharides, which are taken up into the periplasm space. *CjC*hi18A (orange) generates GlcNAc and this lipoprotein may be acting in the periplasmic space (as shown here) or may be located on the outer side of the outer membrane and act outside the cell.

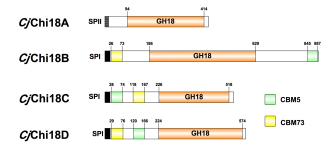


Figure 2. Diverse architecture of the family GH18 chitinases of *C. japonicus.* CAZy domain representation of the family GH18 chitinases of *Cellvibrio japonicus*. The indicated domains are as follows: GH18, family GH18 catalytic domain; CBM5, chitin-binding domain; CBM73, chitin binding domain; SPI, signal peptide, type I; SPII, signal peptide, type II. This image was generated using IBS Illustrator (65).

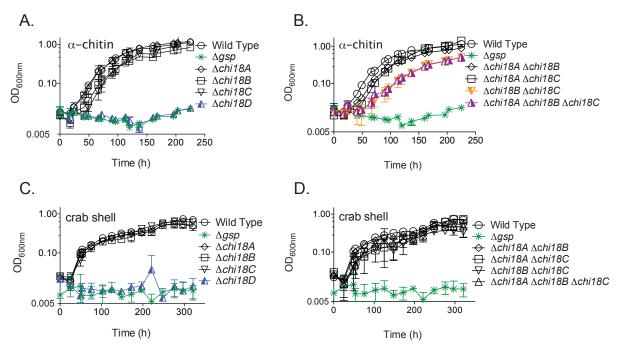


Figure 3. Growth of *C. japonicus* mutants on chitin. Deletion mutants were grown using MOPS minimal medium supplemented with 0.25% α -chitin (A-B) or 4% crab shell (C-D). Panels A and C show single mutants; panels B and D show multiple mutants. All experiments were performed in biological triplicates; error bars represent standard deviations, but in many cases are too small to be observed. These growth experiments were performed simultaneously, but are separated into multiple panels for clarity. As a consequence, the same control strains (wild type and Δgsp) are repeated in each panel for a given substrate.

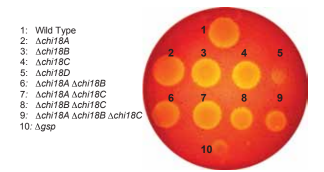


Figure 4. Chitinase secretion of *C. japonicus* GH18 mutants. Strains were grown on a plate that contained MOPS with 1.5% agar, 2% colloidal chitin, and 0.2% glucose. After incubation at 30 °C for 5 days, the plates were stained with Congo Red. This experiment was conducted in biological triplicates, and quantification of the clearing zones is shown in **Table S2**.

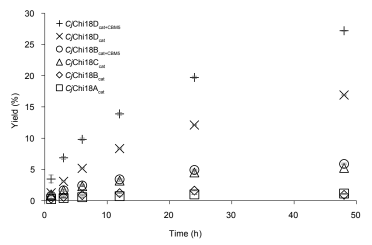


Figure 5. Degradation of α-chitin. Degradation of α-chitin (15 g/L) at 30 °C was tested in 20 mM BisTris pH 6.5, 0.1 mg/mL BSA. The enzyme concentration was 0.5 μ M and samples were taken at different time points. The yield refers to the degree of chitin solubilization. The α-chitin used contained 6.43% ash and 5.42% moisture, and this was taken into account when calculating yields. Each reaction was performed in triplicate; standard deviations are shown as error bars, but are difficult to see since they were low and are partly covered by the symbols.

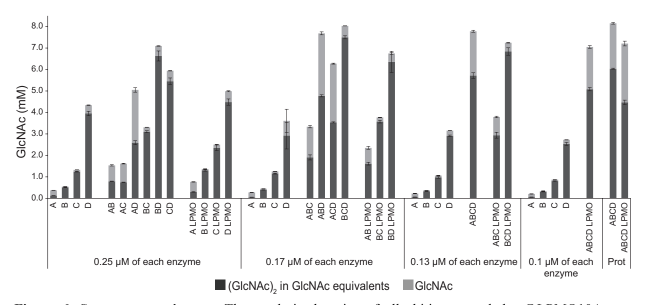


Figure 6. Synergy experiments. The catalytic domains of all chitinases and the *Cj*LPMO10A were mixed in different ways to investigate possible synergistic effects in reactions with α-chitin (15 g/L). The maximum total enzyme load was 0.5 μM with equal amounts of each enzyme, as indicated in the figure. In the reactions where the ratio was determined by protein quantification data from a previous proteomics study ["Prot"; (22)], the total enzyme loading was also 0.5 μM. In the "Prot" reaction without *Cj*LPMO10A the ratio was 21% *Cj*Chi18A, 4% *Cj*Chi18B, 22% *Cj*Chi18C, and 52% *Cj*Chi18D. In the "Prot" reaction with *Cj*LPMO10A the ratio was 14% *Cj*Chi18A, 3% *Cj*Chi18B, 15% *Cj*Chi18C, 35% *Cj*Chi18D, and 33% *Cj*LPMO10A. The *Cj*LPMO10A was Cu²⁺-saturated before use. Reaction mixtures were incubated at 30 °C in 20 mM BisTris pH 6.5, 0.1 mg/mL BSA, and samples were taken after 24 h. In reactions with *Cj*LPMO10A, 0.5 mM ascorbate was added as an external electron donor. Production of GlcNAc and (GlcNAc)₂ was quantified and the amount of (GlcNAc)₂ is given in GlcNAc equivalents. Three parallel reactions were done for each condition and standard deviations are shown as error bars. Reaction mixtures that contained the LPMO showed minor amounts of oxidized (GlcNAc)₂, which were not quantified.

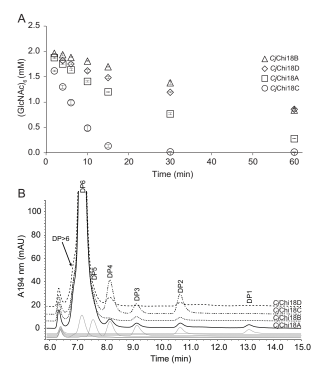


Figure 7. Hydrolysis of (GlcNAc)₆. (A): Hydrolysis of (GlcNAc)₆ over time. The slopes of the linear parts of these curves were used to calculate the initial rates. Standard deviations are shown as error bars. **(B)**: Chromatograms showing the product profile obtained 2 min after mixing chitinases with substrate. DP1-6 represent (GlcNAc)₁₋₆. Chromatograms for various standards are shown as grey lines at the bottom. These experiments were done with the catalytic domains of the chitinases. The reactions contained 2 mM (GlcNAc)₆, 10 mM BisTris pH 6.5, 0.1 mg/mL BSA and 50 nM enzyme, and were done in triplicates at 30 °C. Chromatograms for samples taken after 60 min are shown in **Fig. S4**.

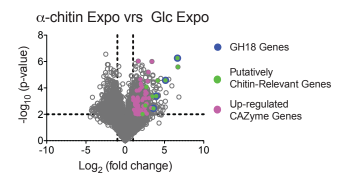


Figure 8. Differential gene expression for *C. japonicus* during exponential growth. This volcano plot shows the $\log_2(\text{fold change})$ plotted against the $-\log_{10}(\text{p-value})$ for every gene in *C. japonicus* during exponential growth on glucose compared to α -chitin, where each gray circle represents the expression of a single gene. The black dashed lines indicate the significance cut-off values: $-\log_{10}(\text{p-value}) > 2$ and $\log_2(\text{fold change}) > 1$. The genes colored magenta represent CAZyme genes, and the complete list of these genes can be found in **Table S3A**.

Systems analysis of the family Glycoside Hydrolase family 18 enzymes from Cellvibrio japonicus characterizes essential chitin degradation functions Estela C. Monge, Tina R. Tuveng, Gustav Vaaje-Kolstad, Vincent G. H. Eijsink and Jeffrey

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