1 Lysozyme activity of the <i>Ruminococcus champanellensis</i>
2 cellulosome
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<ul> <li>Sarah Moraïs<sup>1</sup>, Darrell W. Cockburn<sup>2</sup>, Yonit Ben-David<sup>1</sup>, Nicole M. Koropatkin<sup>2</sup>, Eric</li> <li>C. Martens<sup>2</sup>, Sylvia H. Duncan<sup>3</sup>, Harry J. Flint<sup>3</sup>, Itzhak Mizrahi<sup>4</sup> and Edward A.</li> <li>Bayer<sup>1*</sup></li> <li>9</li> </ul>
10 <sup>1</sup> Biomolecular Sciences Department, The Weizmann Institute of Science, Rehovot,
11 Israel.
12 13 <sup>2</sup> Department of Microbiology and Immunology, University of Michigan Medical
14 School, Ann Arbor, Michigan 48109, USA.
15 16 <sup>3</sup> Microbiology Group, Rowett Institute of Nutrition and Health, University of
17 Aberdeen, Aberdeen, United Kingdom.
<ul> <li>18</li> <li>19 <sup>4</sup> The Department of Life Sciences &amp; the National Institute for Biotechnology in the</li> <li>20 Negev, Ben-Gurion University of the Negev, Beer-Sheva 84105, Israel.</li> <li>21</li> </ul>
22 *Corresponding author:
<ul> <li>Edward A. Bayer, Department of Biological Chemistry, The Weizmann Institute of</li> <li>Science, Rehovot, Israel. Tel: (+972)-8-934-2373. Fax: (+972)-8-934-4118.</li> <li>Email: ed.bayer@weizmann.ac.il</li> </ul>
<ul> <li>26</li> <li>27 Running head: <i>Ruminococcus champanellensis</i> cellulosomal lysozyme</li> <li>28</li> </ul>

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## 29 Originality-Significance Statement

30 This work deals with the identification of a cellulosomal cohesin-containing 31 scaffoldin, which includes a family 25 glycoside hydrolase enzyme that exhibits 32 lysozyme activity. This is the first example of a bacterial lysozyme produced as part 33 of the cellulosome apparatus.

34

# 35 **Abstract**

36 Ruminococcus champanellensis is a keystone species in the human gut that produces 37 an intricate cellulosome system of various architectures. A variety of cellulosomal 38 enzymes have been identified, which exhibit a range of hydrolytic activities on 39 lignocellulosic substrates. We describe herein a unique R. champanellensis scaffoldin, 40 ScaK, which is expressed during growth on cellobiose and comprises a cohesin 41 module and a family 25 glycoside hydrolase (GH25). The GH25 is non-autolytic and 42 exhibits lysozyme-mediated lytic activity against several bacterial species. Despite the 43 narrow acidic pH curve, the enzyme is active along a temperature range from 2 to 44  $85^{\circ}\mathbb{C}$  and is stable at very high temperatures for extended incubation periods. The 45 ScaK cohesin was shown to bind selectively to the dockerin of a monovalent 46 scaffoldin (ScaG), thus enabling formation of a cell-free cellulosome, whereby ScaG 47 interacts with a divalent scaffodin (ScaA) that bears the enzymes either directly or 48 through additional monovalent scaffoldins (ScaC and ScaD). The ScaK cohesin also 49 interacts with the dockerin of a protein comprising multiple Fn3 domains that can 50 potentially promote adhesion to carbohydrates and the bacterial cell surface. A cell-51 free cellulosomal GH25 lysozyme may provide a bacterial strategy to both hydrolyze 52 lignocellulose and repel eventual food competitors and/or cheaters.

53

### 54 Introduction

55

56 Ruminococcus champanellensis is the sole known human gut bacterium able to 57 degrade crystalline cellulose (Chassard et al., 2012). It produces a cellulosome that 58 has been characterized recently (Ben David et al., 2015; Morais et al., 2015). 59 Cellulosomes are large-molecular-weight enzymatic complexes that represent an 60 extremely efficient strategy for cellulose and hemicellulose degradation (Bayer et al., 61 2004; Bayer et al., 2007; Himmel et al., 2010). The strong intermodular calcium-62 dependent cohesin/dockerin interaction drives the assembly between the cellulosomal 63 enzymes and the central non-catalytic, integrating subunit, the scaffoldin, to form the 64 mature complex (Yaron et al., 1995; Lytle et al., 1996). The R. champanellensis 65 genome contains 12 scaffoldins with various molecular arrangements and specificities 66 (Figure 1). The largest cellulosome that could be assembled by *R. champanellensis* 67 would be composed of an anchoring scaffoldin, ScaE, comprising a sortase motif and 68 a cohesin that can interact with the dockerin of a second scaffoldin, ScaB. The latter 69 contains 7 cohesin modules, three of which interact either directly with dockerin-70 bearing enzymes, whereas the remainder can interact either with dockerin-bearing 71 enzymes or with a third dockerin-bearing scaffoldin, ScaA. ScaA also contains two 72 cohesin modules that interact either directly with dockerin-bearing enzymes or 73 indirectly via monovalent scaffoldins (ScaC and ScaD) that play a role of molecular 74 adaptors, thereby modulating the enzymatic composition of the cellulosome (Rincon 75 et al., 2004; Ben David et al., 2015). The entire complex would thus contain a 76 maximum of 11 enzymes. R. champanellensis contains a total of 65 dockerin-bearing 77 proteins, 8 scaffoldin-borne dockerins (Ben David et al., 2015), 25 recently 78 characterized glycoside hydrolases (Morais et al., 2015) and 31 additional dockerin-79 containing proteins. The R. champanellensis cellulosome thus presents a fined-tuned 80 cohesin/dockerin recognition system that enables regulated assembly of an elaborate 81 cellulosomal organization (Morais et al., 2015).

82 Unlike common scaffoldins that are non-enzymatic subunits, the ScaK 83 scaffoldin carries an enzyme that is associated with the metabolism of cellular 84 structural components, i.e., the peptidoglycan. ScaK is composed of a cohesin module 85 (definitive of the scaffoldins) at its N-terminus and a GH25 catalytic domain in its C-86 terminal region. GH25 enzymes are retaining glycoside hydrolases that cleave the  $\beta$ -87 1,4-glycosidic bond between *N*-acetylmuramic acid (NAM) and N-acetylglucosamine

(NAG) in the carbohydrate backbone of bacterial peptidoglycan (i.e., lysozymes).
Two main biological roles can be attributed to bacterial GH25 enzymes: the autolysis,
the re-modeling of peptidoglycan in cellular growth processes, and the dissemination
of phage progeny toward the end of the phage lytic cycle via lysis of the bacterial cell.
These specialized hydrolases can also create enlarged pores in the bacterial
peptidoglycan for the assembly of large trans-envelope complexes (e.g., pili, flagella,
secretion systems) (Vollmer et al., 2008).

95 The genome of *Ruminococcus* sp. CAG:379 strain, closely related to *R*. 96 champanellensis, also contains a gene encoding a protein homologous to ScaK with a 97 similar modular arrangement (cohesin and GH25 modules). In additional 98 *Ruminococcus* species, such as *R. bicirculans* and *R. flavefaciens* that carry multiple 99 GH25 genes, the GH25 module is not accompanied by a cohesin module. ScaK also 100 has 30% identity and 49% similarity with the GH25 module of LytC, an autolysin 101 from *Streptococcus pneumoniae*, which is involved in a three-component mechanism 102 for the lysis of sister cells non-competent for natural genetic transformation (Claverys 103 et al., 2007; Monterroso et al., 2008; Eldholm et al., 2009). The fact that the ScaK 104 GH25 module is connected to a cellulosomal element raises the question of possible 105 lysozyme activity associated with the cellulosome that could provide a novel strategy 106 that would prove beneficial for the bacterial cell.

107 In our previous report (Ben David et al., 2015), the ScaK cohesin was expressed 108 as a CBM-fused cohesin (carbohydrate binding module) but was not found to bind 109 any of the dockerin partners tested. In the present study, we cloned and expressed the 110 ScaK scaffoldin as an intact wild-type protein, in an attempt to reveal potential 111 cohesin affinity partners and to study the GH25 activity.

112 113

#### 114 **Results**

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### 116 Scaffoldin production and functionality of their cohesin modules

117 ScaK, ScaG and ScaF scaffoldins (Figure 1) were produced in *Escherichia* 118 *coli*, purified on Ni-beads, and their estimated molecular weights were in good 119 agreement with their calculated molecular masses (see legend to Supplemental Figure 120 S1). 121 In order to understand the architectural context in which ScaK is positioned 122 within the cellulosome, the dockerin specificities of the ScaK cohesin was examined 123 by an affinity ELISA approach using the representative dockerin-bearing fusion 124 proteins listed in Table 1. For this purpose, 12 dockerin modules were fused to a 125 xylanase tag (xylanase T6 from *Geobacillus stearothermophilus*) that promotes 126 solubility and expression of the dockerin and serves as a recognition tag for the 127 primary antibody in the ELISA procedure (Barak et al., 2005)). The resultant 128 xylanase-dockerin (Xyn-Doc) fusion proteins are listed in Table 1. All dockerins 129 from the R. champanellensis scaffoldins (eight in total) were tested, since they 130 represent the main backbones of its cellulosome structures. The four additional 131 selected dockerins belonged to the 3 dockerin groups (Groups 1, 2 and 3-4) of 132 cohesin-dockerin interactions in this bacterium, as previously defined using 133 bioinformatic-based criteria (Ben David et al., 2015; Morais et al., 2015). ScaK, 134 expressed as a full-length protein (lacking the signal peptide), was able to interact 135 with the ScaG and protein 3939 dockerins (Figure 2A), in contrast to the previous 136 report (Ben David et al., 2015), where the cohesin module was expressed alone, in the 137 absence of the GH25 module.

138 ScaG is a scaffold in that comprises only a cohesin and a dockerin (Figure 1). In 139 a previous report (Ben David et al., 2015), the cohesin and dockerin modules of ScaG 140 were expressed separately, the dockerin demonstrated binding activity for the first 141 cohesin of ScaJ and ScaE, but the cohesin of ScaG failed to interact with any of the 142 dockerin counterparts and was termed inactive in that study. We produced here the 143 full ScaG protein length in an attempt to reveal the binding partner of its cohesin. 144 Using this approach, the ScaG cohesin was able to bind to the ScaA dockerin (Figure 145 1 and Figure 2B), suggesting an adaptor role for this scaffoldin between ScaK and the 146 enzyme-bearing scaffoldin ScaA. A scheme illustrating the interactions between ScaK 147 and the cellulosomal elements is presented in Figure 3. Consequently, the ScaG 148 cohesin is now considered a Group-2 cohesin, and the ScaK cohesin and dockerins of 149 ScaA and Prot3939 belong to the Group-1 interacting modules.

ScaF exhibits the same modular architecture as ScaG (Figure 1), but with a longer linker region at the N-terminus of the protein. Similar to ScaG, when the cohesin and dockerin modules were expressed separately, only the dockerin was active and exhibited similar binding abilities as the ScaG dockerin (Ben David et al., 2015). Similar to ScaK and ScaG, the expression of the full-length scaffoldin revived

the cohesin activity, and specific interaction with the ScaG dockerin was observed (Supplemental Figure S2). As ScaG and ScaF dockerins exhibited the same binding affinities and both can bind directly to the ScaE cohesin, it is not clear why the bacterium would produce and assemble an additional adaptor scaffoldin (i.e., ScaF) to mediate between ScaG and ScaE (via ScaJ) (Supplemental Figure S3). In this context, ScaF could perhaps serve as an extender to increase the overall length of the cellulosome in order to avoid steric hindrance or to access more distant substrates.

162

## 163 Lysozyme activity.

The enzymatic activity was monitored by decrease in turbidity of the cultures. 164 The lysozyme module of ScaK was active on E. coli cells between pH 3.5 and 5.5 165 166 with a pH optimum of 5 (Figure 4A). The activity of the enzyme was lost completely 167 at pH 6 and higher. The protein was stable between 30 and 60°C for 48 h. At 70°C, 168 the lysozyme retained its full activity after 3 h of incubation and was 54% active after 169 24 h of incubation. After 48 h of incubation at this temperature, the enzyme was still 170 30% active. At 80°C, the enzyme was more rapidly degraded, but was still 20% active 171 after 48 h of incubation (Figure 4B). ScaK was active on *E. coli* cells from 2 to 45°C. 172 Enzymatic activity increased with the temperature as monitored by the decrease in 173 turbidity and viable cell count (Figure 4C). In addition, the enzyme was active from 2 174 to 85°C on purified peptidoglycans from *Bacillus subtilis* and *Staphylococcus aureus* 175 along the entire temperature range with moderate enhancement of activity with 176 increasing temperature (Figure 4D).

177 The lytic activity of ScaK was tested against a large number of strains that 178 colonize the human gut and additional bacteria (Table 2). The lysozyme was found to 179 be inactive against the parent R. champanellensis cells but active against several 180 Lactobacillus strains, E. coli, Enterococcus faecalis, Listeria monocytogenes and 181 Streptococcus pneumoniae. Clostridium difficile was degraded after long incubation 182 periods. Growth of E. coli and L. plantarum was inhibited by the presence of the 183 lysozyme, as demonstrated by inhibition zones around disks containing the enzyme 184 during growth on plates (Supplemental Figure S4).

The potential protective role of the bacterial capsule against bacterial cell lysis was tested by incubating the lysozyme with *E. coli* strains producing a capsule (5911 and Nissle strains), and no lytic action was observed. In addition, *Bacteroides thetaiotaomicron*, was tested using both the wild-type strain and a mutant lacking any 189 capsule (Rogers et al., 2013), and in both cases no lysis was observed (Table 2).

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## 191 Expression of ScaK by R. champanellensis.

192 ScaK was detected by proteomic analysis in supernatant fluids of R. 193 *champanellensis* cultures grown on cellobiose as the sole carbon source (score of 9.45 194 and % coverage of 9.23, *p*-value<0.01), suggesting that it is indeed incorporated in the 195 cellulosomal complex and participates in bacterial lignocellulose degradation as a 196 possible inhibitor of other competing or deleterious bacteria which inhabit the human 197 gastrointestinal tract.

198

200

# 199 **Discussion**

The ScaK scaffoldin of *Ruminococcus champanellensis* contains the first cellulosome-associated GH25 known to date. The enzyme is active at extremes of temperature and stable for long incubation periods at elevated temperatures. These characteristics could therefore render the ScaK enzyme a suitable candidate for industrial applications, for example, as an antimicrobial agent in the food industry (Datta et al., 2008; Callewaert et al., 2011; Aminlari et al., 2014) and may also be considered as a therapeutic agent in humans (Pastagia et al., 2013).

208 Curiously, ruminococci seem to carry numerous non-cellulosomal GH25 209 genes - nine in R. flavefaciens FD1, five in R. bicirculans 80/3, five in R. albus 7, one 210 in R. bromii and three in R. champanellensis 18P13 (in addition to the ScaK gene). 211 Since genomic analysis has indicated that prophage DNA (encoding lysozymes) is 212 rather common in this group of bacteria (Berg Miller et al., 2012; Wegmann et al., 213 2014), we analyzed the flanking regions of the ScaK gene for the presence of phage 214 genes, such as holins (Supplemental Table 1). In the absence of such genes, 215 bacteriophage origin would thus be improbable. Therefore, the association of the 216 cohesin module and a GH25 gene only with the R. champanellensis GH25 raises the 217 question of a biological role for this scaffoldin in its cellulosomal complex.

The production of ScaK, ScaG and ScaF as intact proteins revived the binding activity of their lone cohesins towards the specified dockerins, suggesting that the functional status and/or structural stability of the cohesin modules of these scaffoldins is dependent on the full protein architecture. We could thus elucidate their dockerin specificities and therefore complete the proposed architecture of the *R*. *champanellensis* cellulosome system. The results indicate that cell-free cellulosomes can be formed between ScaK that interacts with ScaG, which in turn interacts with ScaA that harbors the enzymes either directly or through ScaC and ScaD adaptors. This type of cellulosome could thus be targeting plant-derived lignocellulosic materials located at a distance from the cell.

228 The ScaK cohesin was also found to interact with the dockerin of protein 229 3939. This very large protein has an estimated molecular weight of 308 kDa, and 230 annotation of its sequence has revealed five predicted Fn3 (fibronectin type III) and 231 two PKD (polycystic kidney disease) domains. The Fn3 domains are relatively 232 common components in celluloytic bacteria, yet their function is not completely 233 understood. Several studies suggested that these domains can mediate protein 234 assembly, adhesion to carbohydrate substrates and/or bacterial cell surfaces (which is 235 in accordance with lysozyme activity). Alternatively, Fn3 domains have been 236 suggested to play a role as flexible peptide linkers or to facilitate the solubility of 237 large protein complexes (Devillard et al., 2004; Alahuhta et al., 2010). In addition, 238 PKD domains have also been found to be involved in protein-protein and/or protein-239 carbohydrate interactions (Lohning et al., 1996). Therefore, we can assume that 240 protein 3939 may have an important, but largely undefined role in carbohydrate 241 degradation, which would benefit the bacterium in general and would justify the 242 expression and secretion of such a large protein. The dockerin of protein 3939 was 243 previously reported to also interact with ScaE that is anchored to the bacterial cell 244 wall (Ben David et al., 2015).

It is also of note that other putative lysozymes can be encoded by GH23-, GH24- and GH73-containing genes in other bacterial genomes. Interestingly, the sole dockerin of the *R. bicirculans* genome was associated with a GH73 catalytic domain (Wegmann et al., 2014), suggesting a similar mechanism and role for the lysozyme in this bacterium.

It is plausible that the GH25 lysozyme has a defensive role against other carbohydrate-degrading bacterial competitors by targeting and effecting the lysis of bacterial cells in the vicinity that would compete for the enzymatic degradation products. In nature, *R. champanellensis* and other specialized fiber-degrading bacteria are prone to exploitation by cheating microbes that utilize the fiber-released soluble sugars, taking advantage of the bacterial fiber-degrading machineries carried without paying the fitness cost (Berlemont and Martiny, 2013). It is therefore reasonable to
assume that the fiber degraders will develop defense mechanisms protecting them
from such exploitation. In this sense, the GH25 lysozyme exhibits an efficient
selective tactic, as it does not affect the parent *R. champanellensis* cell itself, but lyses
other gut microbial cells (Table 2).

261 The bacterial spectrum of susceptibility to ScaK is quite heterogeneous, 262 whereby some Gram-positive (Lactobacillus species, Enterococcus faecalis, Listeria 263 monocytogenes, Streptococcus pneumoniae and Clostridium difficile) and Gram-264 negative (E. coli) bacteria were lysed. Our results initially suggested that ScaK 265 specifically degraded peptidoglycans that contain meso-diaminopimelate (m-DAP) 266 residues in the peptidoglycan structure (Schleifer and Kandler, 1972; Humann and 267 Lenz, 2009), since all the strains that were herein susceptible to ScaK lytic action 268 contained m-DAP. However, since the isolated purified peptidoglycan of S. aureus, 269 which contains D-alanine residue and not m-DAP, was degraded, this hypothesis was 270 thus invalidated.

271 Bacterial defense mechanisms against lysozyme activity are diverse and include 272 the production of enzyme inhibitors or modification of the peptidoglycan (Callewaert 273 et al., 2012), as well as the production of bacterial capsules (Fouet and Mesnage, 274 2002). We thus investigated the role of the bacterial capsule as a possible protection 275 against the lysozyme action by testing *E. coli* strains with and without capsules. In the 276 case of the two encapsulated strains of E. coli, i.e., Nissle and 5911, the capsule 277 indeed conferred resistance to ScaK lytic action. It should also be noted that capsule 278 expression is highly controlled and regulated (Torres-Cabassa and Gottesman, 1987; 279 Gottesman and Stout, 1991; Sledjeski and Gottesman, 1996) and that four different 280 types of bacterial capsules are described to date in *E. coli* (Whitfield and Roberts, 281 1999). The strains tested herein belong to group 2 capsules (capsular gene K5 and 282 K12 for Nissle and 5911 strains, respectively). Therefore, other groups of capsules 283 may confer resistance to the lytic action of ScaK or not, depending also on the growth 284 conditions and extent of capsule expression. We also investigated the capsular 285 protective role with two different strains of Bacteroides thetaiotaomicron (Shah, 286 2013). The two strains tested were the wild type strain and a mutant lacking any 287 capsule. In both cases, no lysis was observed. Consequently, together with the non-288 susceptible Gram-negative bacteria tested herein, the outer membrane may act as an 289 impermeable barrier for small molecules thereby protecting the peptidoglycan layer

from the lytic action of the lysin (Briers and Lavigne, 2015). Nevertheless, some endolysins have been reported to cross the outer membrane and lyse the bacteria (Lai et al., 2011; Lood et al., 2015). This ability could reflect highly positively charged Nor C-terminal domains in their protein sequence, which enable the lysins to bind to the anionic outer membrane and access their peptidoglycan substrate (Lai et al., 2011). In our case, we could not identify a similar domain in ScaK that could account for the lysis of the Gram-negative acapsulated strains of *E. coli* (*BL21* and *5911*).

In conclusion, we identified the presence of a non-autolytic lysozyme in the *R*. *champanellensis* cellulosome system. The production of this cell-free cellulosome would represent a strategy to hydrolyze lignocellulose while repelling eventual food competitors or cheaters. The fact that this lysozyme activity is associated with glycoside hydrolases in a single cell-free cellulosome complex suggests a broader role for cellulosomal complexes that would not be restricted to plant cell wall deconstruction.

304

## 305 **Experimental Procedures**

306 Cloning. Scaffoldins ScaK, ScaG and ScaF were cloned from R. champanellensis 307 genomic DNA using Phusion High Fidelity DNA polymerase F530-S (New England 308 Biolabs, Inc) and the following primers. For ScaK amplification, the primer pair 5'-309 5'ttactaCCATGGcacaccatcaccatcaccatgcagatcagactgtacagac-3' and 310 ttactaCTCGAGttaaaacccattaaatccgt-3' for ScaG, 5'was used. 311 tactgaCCATGGcacaccatcaccatcaccatcagaccatgcagccggcggc-3' 312 5'and 5'-tacttaCTCGAGtcaaccgagcaggtcatccc-3' and for ScaF 313 5'tactgaCCATGGcacaccatcaccatcaccatgcatcggattgacctacag-3' and 314 tacgatCTCGAGtcaccattgcggattcggatc-3'. NcoI and XhoI (restriction sites in 315 uppercase). Fastdigest enzymes (Thermo scientific, USA) were incubated with the 316 PCR products before their ligation into linearized pET28a using T4 DNA ligase (Fermentas UAB, Vilnius, Lithuania). PCR products were purified using a HiYield<sup>TM</sup> 317 318 Gel/PCR Fragments Extraction Kit (Real Biotech Corporation, RBC, Taiwan), and 319 plasmids were extracted using Qiagen miniprep kit (Netherlands). Competent E. coli 320 XL1 cells were used for plasmid transformation.

The full list of fused dockerins used in this article is given in Table 1.

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322 Recombinant protein expression and purification. E. coli BL21 (DE3) cells were 323 transformed with the desired plasmid and plated onto LB-kanamycin plates. The cells 324 producing ScaK, ScaG and ScaF were grown in 500 ml LB (Luria Broth) and 2 mM 325 CaCl<sub>2</sub> at 37°C until  $A_{600} \approx 0.8$ -1. The cells were induced by adding 0.1 mM (final 326 concentration) of isopropyl-1-thio-β-D-galactoside (IPTG) (Fermentas UAB Vilnius, 327 Lithuania), and cell growth was continued at 16°C overnight. Cells were harvested by 328 centrifugation at 5000 rpm for 5 min. Pelleted cells were resuspended in 1 ml TBS 329 (Tris-buffered saline, 137 mM NaCl, 2.7 mM KCL, 25 mM Tris-HCl, pH 7.4). Xyn-330 Doc proteins were expressed as described previously (Ben David et al., 2015; Morais 331 et al., 2015).

His-tagged proteins (scaffoldins and Xyn-Doc) were purified on a Ni-NTA
column (Qiagen), as reported earlier (Caspi et al., 2006).

Purity of the recombinant proteins was tested by SDS-PAGE on 10% acrylamide gels, and protein concentration was estimated by absorbance (280 nm), based on the known amino acid composition of the protein, using the Protparam tool (Gasteiger et al., 2005). Proteins were stored in 50% (v/v) glycerol at -20°C.

338 Affinity-based ELISA. The matching fusion-protein procedure of Barak et al (Barak 339 et al., 2005; Caspi et al., 2006) was followed to determine cohesin-dockerin 340 specificity of interaction. Scaffoldins were immobilized on the plate at a concentration 341 of 1  $\mu$ g/ ml (100  $\mu$ l/well) in 0.1 M sodium carbonate (pH 9) and incubated at 4°C 342 overnight. The following steps were performed at room temperature for 1 h with all 343 reagents at a volume of 100  $\mu$ /well with a washing step (300  $\mu$ /well blocking buffer 344 without BSA) repeated three times after each step. The coating solution was 345 discarded, and blocking buffer (TBS, 10 mM CaCl<sub>2</sub>, 0.05% Tween 20, 2% BSA) was 346 added. The blocking buffer was discarded, and the desired representative Xyn-Doc(s) 347 (Xyn-DocA, Xyn-DocB, Xyn-DocC, Xyn-DocD, Xyn-DocF, Xyn-DocG, Xyn-DocH, 348 Xyn-DocJ, Xyn-Doc3939, Xyn-Doc48, Xyn-Doc9b, Xyn-Doc3550), diluted to 349 concentrations of 0.1, 1, 10 and 100 ng/ml in blocking buffer, were added. Rabbit 350 anti-xylanase antibody (diluted 1:10000) was used as the primary antibody 351 preparation, and secondary antibody preparation was HRP-labeled anti-rabbit 352 antibody diluted 1:10000 in blocking buffer. Substrate-Chromogen TMB (Dako, 353 Agilent Technologies, USA) was added at 100 µl/well, and the reaction was carried 354 out for 2 min before color formation was terminated upon addition of 1 M  $H_2SO_4$  (50

 $\mu$ /well), and the absorbance was measured at 450 nm using a tunable microplate reader.

357 **Enzymatic activity assay.** All assays were performed at least twice in triplicate. 358 ScaK enzymatic activity was monitored using the turbidimetric method (Shugar, 359 1952; Diez-Martinez et al., 2015) with modifications. The enzyme was applied at a concentration of 50 µg/ml, and E. coli BL21 at OD<sub>600nm</sub>= 1 was used as a substrate 360 361 for examining pH and temperature effects on the enzyme. After 30 min incubation 362 time, decreases in optical densities were read at 600 nm after centrifuging 2 min at 363 1000 rpm (60  $\times$  g) to clear cell debris. At this centrifugal speed, less than 10% of the 364 bacterial cells precipitate, for short time periods (2 min) as can be observed in 365 Supplemental Figure S5A and S5B. Without the centrifugation step, cell debris (that 366 precipitate only gradually over extended time periods) can float and OD 367 measurements can be less accurate. Controls without enzyme were also submitted to 368 the centrifugal step so that the 10% precipitation of the bacterial cells does not affect 369 the results. Supplemental Figure S5C presents the effect of temperature on lytic 370 activity after 1 h incubation without the centrifugation step, and the results tend to be 371 similar to those presented in Figure 4C with the centrifugation step.

372 The pH optimum was determined by using acetate buffer ranging from pH 3.5 373 to 6.5 and MOPS (3-(N-morpholino)propanesulfonic acid) from pH 6.5 to 9. The 374 enzyme was incubated at 37°C for 15 min. Temperature optima were tested at pH 5 375 (50 mM acetate buffer) between temperatures ranging from 2 to 45°C, and reactions 376 were terminated after 30-min incubation. We used a viability assay (Diez-Martinez et 377 al., 2015) to follow the temperature effect on ScaK lytic action. Measurement of 378 viable *E. coli* cells was carried out in LB agar plates after 30-min incubation at each 379 temperature. For each sample, a 10-fold dilution series was prepared in LB, and 100 380 ul of each dilution was plated. Colonies were counted after overnight incubation at 37°C. 381

The susceptibility of the peptidoglycans from *B. subtilis* and *S. aureus* (Sigma-Aldrich, Israel) to lysozyme was analyzed with a turbidometric assay (Bera et al., 2005; Wang et al., 2009). The peptidoglycans were diluted to 0.5 mg/ml in double distilled water, and the lysozyme was added to a concentration of 50  $\mu$ g/ ml at pH 5 (50 mM acetate buffer). The absorbance at 450 nm was monitored after 30 min of

Page 13 of 27

incubation at temperatures ranging from 2 to 85°C.

Stability of the protein was tested for 30 min at 37°C after incubation for 1, 3,
24 and 48 h at 30, 40, 50, 60, 70 and 80°C.

390 Bacterial strains and growth medium. The bacterial species tested in this study are 391 listed in Table 2. All the strains were cultured at 37°C (anaerobic strains were grown 392 under anaerobic conditions) and tested for their sensitivity to ScaK in vitro as 393 described above (at 1 h incubation time). Certain strains listed in Table 2 were held in 394 stock by the authors S.H. Duncan and H.J. Flint (U. of Aberdeen). The 18 cultures 395 provided for testing for ScaK lytic activity were Bifidobacterium adolescentis L2-32, 396 Collinsella aerofaciens DSM 3979, Enterococcus faecalis JH2-2, Eubacterium hallii 397 L2-7 (DSM 17630), Anaerostipes hadrus Ss2/1, Blautia obeum A2-162, Eubacterium 398 *rectale* A1-86 (DSM 17629), Roseburia faecis M72/1 (DSM 16840), 399 Lachnospiraceae sp. nov. M62/1, Lactobacillus reuteri Ca6, Ruminococcus 400 18P13 18848), champanellensis (DSM Ruminococcus bicirculans 80/3. 401 Ruminococcus bromii L2-63, Eubacterium siraeum 70/3, Faecalibacterium 402 prausnitzii A2-165 (DSM 17677), Ruminococcus flavefaciens 17, Ruminococcus 403 albus SY3 and Streptococcus gordonii DL-1. Those available from Deutsche 404 Sammlung von Mikroorganismen und Zellkulturen (DSMZ), Germany) are indicated 405 by DSM numbers in brackets. Cultures were prepared by growing on M2GSC 406 medium (Miyazaki et al., 1997) for approximately 24 h under CO<sub>2</sub> prior to testing for 407 ScaK lysis. E. coli BL21 and Nissle, Bacillus subtilis NCIB 3610 and PY79 strains, 408 Acinetobacter baumanii ATCC 17978, Listeria monocytogenes 10403S and Vibrio 409 *cholerae* TRH7000 were cultured in LB. Bacteroides thetaiotaomicron ATCC 29148 410 and  $\Delta CPS$  (Cameron et al., 2014) were cultured in TYG (Holdeman et al., 1977). 411 Lactobacillus plantarum WCFS1, Lactobacillus pentosus DSM 20314, Lactobacillus 412 reuteri Ca6 and Lactobacillus rhamnosus GG were cultured in MRS. Lactococcus 413 lactis MG5267 and Z3000 strains were cultured in M17. Clostridium difficile VPI 414 10463 was cultured in BHIS (Sorg and Dineen, 2009). Clostridium thermocellum 415 DSM 1313 and Clostridium clarifavum DSM 19732 were cultured in GS-2 medium. 416 Ruminococcus champanellensis 18P13 was cultured in M2 medium. Streptococcus 417 pneumoniae R6 was grown in Todd Hewitt broth supplemented with 5% yeast extract 418 (Updyke and Nickle, 1954).

419

E. coli strain 5911 was grown for 3 h in LB, and capsule expression was

420 induced by addition of 15% sucrose (Sledjeski and Gottesman, 1996). The cells were

- 421 then grown overnight at 37°C and incubated at room temperature for a week.
- 422 Bacterial cells were tested by the turbidimetric assay as described above.

Inhibition test. *E. coli* BL21 and *L. plantarum* WCFS1 cells were spread on the entire
surface of LB or MRS plates prepared at pH 5. Sterile filter paper disks (catalogue
number 74146, Sigma Aldrich, Israel) containing either sterile water or 2 g/l ScaK
were placed in the middle of the plates, and the plates were incubated overnight at
37°C.

428 Label free LC MS-MS analysis. Proteolysis and mass spectrophometry analysis of
429 *R. champanellensis* culture supernatants, grown on cellobiose as a carbon source
430 (Lopez-Siles et al., 2012; Morais et al., 2015), were performed as described by Artzi
431 (Artzi et al., 2015).

432

## 433 Acknowledgements

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445

### 446 **Conflict of Interest**

- 447 The authors declare no conflict of interest
- 448

Table 1: List of the *R. champanellensis* Xyn-fused dockerin proteins used in this
study. Name and modular architecture of the original scaffoldin or protein are given.
Abbreviations: Xyn, XynT6 from the *G. stearothermophilus*; SIGN, signal peptide;
Doc, dockerin; Coh, cohesin; GH, glycoside hydrolase; SGNH, lipases or esterases;
Fn3, fibronectin type III; PKD, polycystic kidney disease; UNK, X, unknown.

454 The numbers for proteins 3550 and 3939 refer to the last four digits of the respective

455 full GI number (i.e. 29154XXXX).

456

Fused dockerin		Parent	Modular architecture	Dockerin group	
		scaffoldin/protein			
	Xyn-DocA	ScaA	SIGN X Coh Coh <b>Doc</b>	2	
	Xyn-DocB	ScaB	SIGN Coh Coh Coh Coh Coh	1	
			Coh Coh <u>X</u> <b>Doc</b>		
	Xyn-DocC	ScaC	SIGN Coh UNK <b>Doc</b>	2	
	Xyn-DocD	ScaD	SIGN Coh <b>Doc</b>	2	
	Xyn-DocF	ScaF	SIGN Coh <b>Doc</b>	1	
	Xyn-DocG	ScaG	SIGN Coh <b>Doc</b>	1	
	Xyn-DocH	ScaH	SIGN SGNH Coh <b>Doc</b>	1	
	Xyn-DocJ	ScaJ	SIGN Coh Coh Coh <mark>Doc</mark>	1	
	Xyn-Doc3939	gi:29154 <u>3939</u>	SIGN Fn3 PKD Fn3 Fn3 Fn3	1	
9			Fn3 PKD <b>Doc</b>		
4	Xyn-Doc48	Cel48A	SIGN GH48A <b>Doc</b>	2	
	Xyn-Doc9B	GH9B	SIGN CBM4 Fn3 GH9B Doc	3-4	
			GH16A		
- 1	Xyn-Doc3550	gi:29154 <u>3550</u>	UNK_ <b>Doc</b>	3-4	

457

459

- 460 Table 2: Bacterial strains tested for ScaK lytic action. Enzymatic activity was monitored by decrease in turbidimetry in cell cultures at 600 nm, from the beginning 461 462  $OD_{T=0}$  to the end of the reaction  $OD_{T=1h}$ . Classification of the Firmicutes based on Ludwig et al (Ludwig et al., 2009). 463
- 464 <sup>a</sup> grown with sucrose

465

	Bacterial phyla/	Gram	OD <sub>T=0</sub>	OD <sub>T=1h</sub>	ScaK lytic action
	(family)/species/strain		10	1 10	J.
	Actinobacteria				
	(Bifidobacteriaceae)				
	Bifidobacterium adolescentis L2-32	+	0.132	0.124	_
	(Coriobacteriaceae)				
	Collinsella aerofaciens DSM 3979	+	0.094	0.095	_
	Firmicutes				
	(Bacillacae)				
	Bacillus subtilis NCIB 3610	+	1.200	1.195	_
	Bacillus subtilis PY79	+	1.220	1.199	-
- (					
	(Enterococcaceae)				
	Enterococcus faecalis JH2-2	+	0.299	0.210	+
	(Lachnospiraceae)				
	Eubacterium hallii L2-7	+	0.085	0.074	_
	Eubacterium rectale A1-86	+	0.925	0.884	-
	(DSM17629)				
	Anaerostipes hadrus SS2/1	+	1.100	1.029	_
	Lachnospiraceae sp. nov. M62/1	+	0.085	0.088	_
	Blautia obeum A2-162	+	1.030	0.984	-
	Roseburia faecis M72/1	+	0.862	0.856	_
	(Lactobacillaceae)				
	Lactobacillus plantarum WCFS1	+	1.840	0.368	+
	Lactobacillus pentosus DSM 20314	+	1.560	0.267	+
	Lactobacillus reuteri Ca6	+	0.210	0.214	_
	Lactobacillus rhamnosus GG	+	1.952	0.341	+
	(Listeriaceae)				
	Listeria monocytogenes 10403S	+	1.194	0.467	+
	(Peptostreptococcaceae)				
	Clostridium difficile VPI 10463	+	0.470	0.418	±
				(T=4h)	
	(Ruminococcaceae)				
	Ruminococcus champanellensis 18P13	+	1.130	1.259	-

	Ruminococcus bicirculans 80/3	+	0.459	0.444	_
	Ruminococcus bromii L2-63	+	0.382	0.361	_
	Eubacterium siraeum 70/3	+	0.437	0.422	_
	Faecalibacterium prausnitzii A2-165	+	0.336	0.335	_
	Ruminococcus flavefaciens 17	+	0.331	0.330	_
	Ruminococcus albus SY3	+	0.689	0.665	_
	Clostridium clariflavum DSM 19732	+	1.345	1.322	_
	Clostridium thermocellum DSM 1313	+	1.202	1.197	_
	(Streptococcaceae)				
	Lactococcus lactis MG5267	+	1.500	1.501	_
	Lactococcus lactis Z3000	+	1.500	1.499	_
	Streptococcus gordonii DL-1	+	0.109	0.100	_
	Streptococcus pneumoniae R6	+	0.480	0.051	+
	* *				
	Bacteroidetes				
	(Bacteroideaceae)				
	Bacteroides thetaiotaomicron ATCC	_	1.573	1.559	_
	29148				
	Bacteroides thetaiotaomicron $\Delta CPS$	_	1.567	1.509	_
	Proteobacteria				
	(Enterobacteriaceae)				
	Escherichia coli BL21	-	1.462	0.206	+
	Escherichia coli 5911 (K12)	-	1.500	0.190	+
	Escherichia coli 5911 (K12) <sup>a</sup>	_	1.423	1.360	_
	<i>Escherichia coli</i> Nissle	_	1.692	1.666	_
	(Moraxellaceae)				
	Acinetobacter baumanii ATCC 17978	_	0.689	0.648	_
	(Vibrionaceae)				
	Vibrio cholera TRH7000	_	0.656	0.619	_
466					
100					

# 467 **Figure legends**

468

**Figure 1:** Schematic representation of the cohesin-bearing scaffoldin proteins in *R. champanellensis* based on the respective genome sequences. SGNH, hydrolase-type esterase domain (IPR013830); GH25, a putative GH25-family module sharing similarity to lysozyme. Specific interactions according to sequence alignment and biochemically characterized cohesin/dockerin interactions are color-coded. Shaded colors indicate that the designated cohesins or dockerins exhibit more selective interactions with only some of their Group 1 or Group 2 counterparts.

- 476
- 477

Figure 2: Newly discovered dockerin-binding profiles of R. champanellensis ScaK 478 479 and ScaG cohesins measured by affinity ELISA (A) ELISA experiments 480 demonstrating different interaction specificities between the ScaK cohesin and 481 selected scaffoldin- and enzyme-borne dockerins. ScaK interacted with DocG and 482 Doc3939 (B) ELISA experiments demonstrating different interaction specificities 483 between the ScaG cohesin and selected scaffoldin- and enzyme-borne dockerins. 484 ScaG interacted exclusively with DocA. Error bars indicate the standard deviation 485 from the mean of triplicate samples (ELISA). The experiments were performed three 486 times.

487

488

Figure 3: Proposed cell-free cellulosome complexes involving ScaK in *R. champanellensis*. Different types of cohesin-dockerin interactions are color-coded.
The ScaK cohesin binds selectively to the dockerins of ScaG and Prot3939 and not to other Group 1 dockerins. Similarly, the ScaG cohesin appears to be very selective in its binding to the ScaA dockerin, and fails to bind other Group 2 dockerins.

494

495 Figure 4: Characterization of recombinant R. champanellensis ScaK. (A) Effect of 496 pH on lysozyme activity after 15 min incubation. (B) Thermostability of ScaK-497 derived lysozyme activity at different temperatures, following the heat-shock, the 498 enzymatic activity was measured at pH 5 after 30 min incubation at 37°C. (C) Effect 499 of temperature on lysozyme activity on *E. coli* cells at pH 5 after 30 min incubation, 500 solid line representing the decrease in OD at 600 nm and dashed line, the viability of the cells (in cfu/ml) with the control counting 1.6.10<sup>12</sup> cfu/ml. (D) Effect of 501 502 temperature on lysozyme activity on the peptidoglycans of B. subtilis (blue line) and 503 S. aureus (red line) at pH 5 after 30 min incubation. Enzymatic reactions, performed 504 using the modified turbidimetric method were repeated in triplicate, and standard 505 deviations are indicated.

506

## 507 Supplemental Figures

508

Supplemental Figure S1: Purity of the recombinant scaffoldins after Ni-NTA
purification as assessed by SDS-PAGE gels. A. ScaK, molecular weight 52326 Da
(10% acrylamide), B. ScaG, molecular weight 29510 Da (12% acrylamide) and C.
ScaF, molecular weight 25633 Da (12% acrylamide).

513

514 **Supplemental Figure S2:** *R. champanellensis* ScaF interactions measured by ELISA 515 with selected dockerins. The cohesin only binds to the dockerin of ScaG. Error bars 516 indicate the standard deviation from the mean of triplicate (ELISA) from three 517 experiments.

518

**Supplemental Figure S3:** Proposed cell-bound cellulosome complexes involving ScaG and ScaF in *R. champanellensis*. Different types of cohesin-dockerin interactions are color-coded. The ScaG and ScaF dockerins are selective for the cohesins of ScaE and ScaJ1. The ScaG dockerin also binds selectively to the ScaF cohesin, but the ScaF cohesin fails to bind to its own dockerin. The ScaG cohesin appears to be very selective in its binding to the ScaA dockerin and fails to bind other Group 2 dockerins.

526

527 Supplemental Figure S4: Inhibition of *L. plantarum* (A) and *E. coli* (B) growth by
528 the presence of the GH25 lysozyme on disks (2 g/l). The cells were grown on MRS
529 and LB plates prepared at pH 5. Inhibition zones are marked with red arrows.

530

531 Supplemental Figure S5: Effect of time on the precipitation of bacterial cells *L*.
532 *plantarum* (A) and *E. coli* (B) at a centrifugal speed of 1000 rpm (60 x g). (C) Effect
533 of temperature on lysozyme activity on *E. coli* cells after 1 h incubation at pH 5
534 without the centrifugation step.
535

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537

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Figure 1: Schematic representation of the cohesin-bearing scaffoldin proteins in R. champanellensis based on the respective genome sequences. SGNH, hydrolase-type esterase domain (IPR013830); GH25, a putative GH25-family module sharing similarity to lysozyme. Specific interactions according to sequence alignment and biochemically characterized cohesin/dockerin interactions are color-coded. Shaded colors indicate that the designated cohesins or dockerins exhibit more selective interactions with only some of their Group 1 or Group 2 counterparts. 80x94mm (300 x 300 DPI)



Figure 2: Newly discovered dockerin-binding profiles of R. champanellensis ScaK and ScaG cohesins measured by affinity ELISA (A) ELISA experiments demonstrating different interaction specificities between the ScaK cohesin and selected scaffoldin- and enzyme-borne dockerins. ScaK interacted with DocG and Doc3939 (B) ELISA experiments demonstrating different interaction specificities between the ScaG cohesin and selected scaffoldin- and enzyme-borne dockerins. ScaG interacted exclusively with DocA. Error bars indicate the standard deviation from the mean of triplicate samples (ELISA). The experiments were performed three times.

109x40mm (300 x 300 DPI)

Accepted



Figure 3: Proposed cell-free cellulosome complexes involving ScaK in R. champanellensis. Different types of cohesin-dockerin interactions are color-coded. The ScaK cohesin binds selectively to the dockerins of ScaG and Prot3939 and not to other Group 1 dockerins. Similarly, the ScaG cohesin appears to be very selective in its binding to the ScaA dockerin, and fails to bind other Group 2 dockerins.

109x65mm (300 x 300 DPI)

Accept



Figure 4: Characterization of recombinant R. champanellensis ScaK. (A) Effect of pH on lysozyme activity after 15 min incubation. (B) Thermostability of ScaK-derived lysozyme activity at different temperatures, following the heat-shock, the enzymatic activity was measured at pH 5 after 30 min incubation at 37°C. (C) Effect of temperature on lysozyme activity on E. coli cells at pH 5 after 30 min incubation, solid line representing the decrease in OD at 600 nm and dashed line, the viability of the cells (in cfu/ml) with the control counting 1.6.1012 cfu/ml. (D) Effect of temperature on lysozyme activity on the peptidoglycans of B. subtilis (blue line) and S. aureus (red line) at pH 5 after 30 min incubation. Enzymatic reactions, performed using the modified turbidimetric method were repeated in triplicate, and standard deviations are indicated. 109x71mm (300 x 300 DPI)

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