In previous work we have described *Fasciola gigantica* DM9-1 (FgDM9-1) as a 16.8 kDa cytoplasmic protein which showed intracellular relocalization to vesicular structures following bacterial, drug and heat stress [1]. This was similar to the behavior observed for *Anopheles gambiae* PRS1 (AgPRS1) which relocated from the cytoplasm to vesicles upon infection with *Plasmodium* [2]. AgPRS1 was detected in salivary glands and midgut whereas FgDM9-1 was present in parenchymal cells. Both proteins carried a pair of DM9 domains that were described in protein family databases as domains of unknown function, e.g., IPR006616 (InterPro), SM00696 (Smart), DUF3421 (Pfam), and 128937 (CDD). Furthermore, recombinant FgDM9-1 showed a tendency to oligomerize and the protein was predicted to have a β-strand rich secondary structure. The same behavior was observed for *Opisthorchis viverrini* DM9-1 a homologous (but not orthologous) protein with 2 DM9 domains that was detected in the tegumental layer [3]. Circular dichroism analysis of OvDM9-1 supported a high β-strand content of DM9 proteins. Structure analysis in Phyre2 showed similarity to *Ascaris* cytoplasmic motility protein (MFP2), a protein found in sperms that contains 4 DM9 domains [4]. A biochemical activity could not be assigned, neither to MFP2 nor to PRS1, FgDM9-1, and OvDM9-1. Shortly afterwards *Crassostrea gigas* lectin 1 (CGL1), a homologous 15.5 kDa protein with 2 DM9 domains, was described as a novel lectin with high mannose specificity [5,6]. CGL1 contained 2 mannose-binding sites that were unrelated to any previously described mannose-binding site. Each of the 2 DM9 domains contributes residues to each binding site and some of these residues show high conservation among proteins with DM9 domains.

This study investigated whether FgDM9-1 was capable to bind mannose as CGL1 and if this lectin activity extended to bacteria and human erythrocytes with mannosylated molecules on their surface. Concanavalin A (ConA), an extensively researched protein of the jack bean with comparable lectin properties [7,8] but a distinct mannose-binding site [9] was used for comparison. The structural conservation of the mannose-binding sites between CGL1 and FgDM9-1 was theoretically analyzed to understand how residues can be substituted without disturbing binding activity.

The resolved tertiary structure of CGL1 [5] (PDB: 5IDB) was used as a template to build a model of FgDM9-1 in SWISS-MODEL [10]. The amino acid alignment created during the modelling process is shown in Fig. 1A. Sequence conservation between the 2 proteins is modest with 42 residues fully conserved resulting in an identity of 26.2% as calculated in EM-
BOSS aligncyppair [11]. FgDM9-1 carries ten additional residues at the N-terminus and one additional residue at the C-terminus. Larger internal differences are 2 indels, a 5-residue deletion and one additional residue at the C-terminus. Larger internal differences are 2 indels, a 5-residue deletion and a 5-residue insertion. The 2 mannose-binding sites identified in CGL1 [5, 6] show high conservation in FgDM9-1 with the exception of an aspartic acid (Asp22) in the first binding site (Fig. 1B). This residue is missing due to the 5-residue deletion in FgDM9-1 and while it seems to be essential for in vitro mannose-binding of CGL1 [6] there is at present no proof that this is also the case in vivo. In any way, the second binding site of CGL1 contains only 5 residues lacking an equivalent aspartic acid. On the other hand, the 5-residue insertion carried by FgDM9-1 and other trematode DM9 proteins might provide a residue that contributes to mannose-binding in binding site 2. The 5 residues include an aspartic acid, and this residue is also present in Anopheles gambiae PRS1, a homologous protein with 2 DM9 domains [2]. In the model none of these 5 residues is close enough to interact with mannose in the second binding site. This might be a limitation of the automated calculation because there is no corresponding...
CGL1 sequence to model on. In addition to the lack of aspartic acid in binding site 1 FgDM9-1 carries a glycine in place of alanine (CGL1: DKHEFA, FgDM9-1: -KHEFG) and binding site 2 has 2 substitutions tyrosine for phenylalanine, glycine for alanine (CGL1: KYEFA, FgDM9-1: KYEYG). In both proteins histidine in binding site 1 is substituted by tyrosine in binding site 2. These substitutions are conservative because the involved amino acids are similar in their physicochemical properties and should support the binding of mannose. The always found lysine and glutamic acid in the binding sites are also highly conserved in the DM9 domains of other proteins suggesting that they are the most important residues in this novel mannose-binding site. Substitution of lysine in binding site 1 by alanine knocked out mannose-binding in CGL1 [6].

The complete 468 nt open reading frame of the FgDM9-1 transcript (GenBank: KT865082) [1] was amplified by PCR. The primers 5’-cat atg GCG CAA TGT GGT CAC G-3’ and 5’-ctc gag TCA TTT CTT CAT CCA CAC-3’ introduced terminal Nde 1 and Xho I restriction endonucleases sites (lower case) at start- and stop-codon (underlined), respectively. These sites were used to insert the DNA fragment into the expression vector pET21b(+). Recombinant FgDM9-1 without His-tag was expressed in Escherichia coli BL-21(DE3) and purified from the soluble bacterial protein fraction by affinity chromatography on D-mannose agarose. Mannose in the eluate was depleted by filtration and washes on 10 kDa cutoff centrifugal filter devices. The concentrated protein solution was transferred to a microtube and kept at −20°C. The plasmid pQE30-FgDM9-1 created in a previous analysis was used to obtain recombinant FgDM9-1 with an N-terminal His-tag as described [1]. The Bio-Rad Protein Assay was used to determine protein concentration. Only recombinant FgDM9-1 expressed without His-tag showed mannose-binding and migrated as a monomer at the expected molecular mass in SDS-PAGE (Fig. 2). SDS-PAGE showing the purification of E. coli expressed soluble FgDM9-1 by mannose affinity chromatography. M: Broad range protein standard marker (Bio-Rad, Hercules, California USA). Lane 1: flow through, lanes 2-3: wash fractions, lanes 4-7: elution fractions. Recombinant FgDM9-1 migrates as a monomer at the expected molecular mass of 16.8 kDa. The result strongly supports the structure model of FgDM9-1 built on CGL1 with 2 conserved mannose-binding sites. It also suggests that aspartic acid as found in CGL1 binding site 1 is not essential for binding of mannose by FgDM9-1.

Forty EDTA red blood cell (RBC) samples classified by their ABO group from Blood Bank Thammasat University Hospital and stored at 4°C until use. The protocol was reviewed and approved by the Ethical Review Sub-Committee Board for Human Research Involving Sciences of Thammasat University (COE no. 007/2562). Hemagglutination activity of FgDM9-1 was assayed on a microtiter plate following Adamová and colleagues [12]. RBC were packed by centrifugation, washed in PBS, packed again and resuspended in PBS to a final concentration of 1% v/v. A volume of 50 µl of FgDM9-1 [1 mg/ml] was pipetted into each well of a 96-well FASTEC U plate. ConA at the same concentration was used as a positive control and PBS was used as a negative control. In the next step 50 µl of 1% RBC were added to each well, mixed and incubated at room temperature for 45 min until the negative control had fully sedimented. Agglutination reactions were visually observed and graded as positive/negative. These experiments were done to evaluate whether FgDM9-1 was able to cross-link cells. Such activity is only possible if a single protein contains either multiple binding sites or if it oligomerizes to present more than one binding site. All samples regardless of blood group showed agglutination following incubation with either FgDM9-1 or ConA (Fig. 3).

Twelve gram-positive and -negative bacterial strains (Table 1) were kindly provided by Assistant Professor Dr. Potjanee Sriamanote (Microbiology, Graduate Program in Biomedical Sciences, Faculty of Allied Health Sciences, Thammasat University). The protocol was reviewed and approved by the Biosafety Committee of Thammasat University (Ref. No. 065/2562). All bacteria were grown in LB medium at 37°C, 250 rpm shaking until an OD600 of 0.5. A volume of 50 µl of each bacterial suspension was pipetted into the wells of a 96-well FASTEC U plate. Then 50 µl of FgDM9-1 in TBS [1 mg/ml] were added and incubated at 37°C, 16 hr. ConA was used as a positive control at the same concentration. Agglutination reactions were visually observed and graded as positive/negative. Streptococcus group A was used to find the minimum amount of
Table 1. Bacterial agglutination

| Bacterial strain                        | Agglutination |          |          |          |          |          |          |          |          |
|----------------------------------------|---------------|----------|----------|----------|----------|----------|----------|----------|----------|
|                                        | ConA          | FgDM9-1  | ConA     | FgDM9-1  | ConA     | FgDM9-1  | ConA     | FgDM9-1  | ConA     |
| *E. coli* Top10                         | Gram-negative rod | +       | +        |          |          |          |          |          |          |
| *Shigella* group D                      | Gram-negative rod | +       | +        |          |          |          |          |          |          |
| Methicillin-resistant *Staphylococcus aureus* (MRSA) | Gram-positive cocci  | +       | +        |          |          |          |          |          |          |
| *Streptococcus* group A                 | Gram-positive cocci  | +       | +        |          |          |          |          |          |          |
| *Streptococcus pneumoniae*              | Gram-positive lance-shaped cocci | +       | +        |          |          |          |          |          |          |
| *Vibrio cholerae* VP03:k6               | Gram-negative curved rods | +       | +        |          |          |          |          |          |          |
| *Serratia marcescens*                   | Gram-negative rod | −       | −        |          |          |          |          |          |          |
| *E. coli* ETEC LT+ ST-                 | Gram-negative rod | −       | −        |          |          |          |          |          |          |
| *E. coli* AAEC TH                       | Gram-negative rod | −       | −        |          |          |          |          |          |          |
| *Enterococcus* spp.                    | Gram-positive cocci | −       | −        |          |          |          |          |          |          |
| *Yersinia enterocolitica*               | Gram-negative coccolbacillus | −       | −        |          |          |          |          |          |          |

FgDM9-1 that caused agglutination. FgDM9-1 protein solution was 2-fold diluted from 1 mg/ml to $4.88 \times 10^{-4}$ mg/ml and tested as above against 50 µl of bacterial suspension. All assays were performed 2 times with independently prepared FgDM9-1. Many but not all bacterial species carry mannose containing glycosylation on their surface. Six of the twelve tested bacterial strains were agglutinated by FgDM9-1 and ConA (Table 1). Only a single species, *Serratia marcescens*, was agglutinated by FgDM9-1 but not ConA. Five bacterial samples were neither agglutinated by FgDM9-1 nor ConA. The minimum concentration of FgDM9-1 that could fully agglutinate *Streptococcus* group A was 62.5 µg/ml (Fig. 4).

In conclusion, FgDM9-1 showed mannose-binding and lectin activity that facilitated agglutination of pro- and eukaryotic cells similar to ConA. The plant lectin ConA is a homотetrameric protein formed by 26.5 kDa subunits, each with one mannose/glucose-binding site [9]. Monomeric FgDM9-1 contains 2 potential mannose-binding sites comparable to the homologous CGL1. The 2 studies concerning CGL1 in the oyster *Crassostrea gigas* showed occupation of both binding sites by mannose in the crystallized CGL1 but found only the first site comprising 6 residues occupied by mannose in in vi-
tro studies or upon isolation of the native protein [5,6]. The second site lacks an aspartic acid residue which was argued by the authors to stabilize mannose-binding through additional bonds. Substitution of residues in the first binding site of CGL1 (D22A, K43A, H52A) supported the importance of these residues in binding [6]. Unfortunately, these mutant forms were not tested in presence of mannose in crystallization analysis to confirm the results in a situation that demonstrated binding of 2 mannose residues of the wild type sequence. Of course, a simple in vivo assay might not be able to mimic the acute intracellular situation in which the protein binds mannose. What are these situations? PRS1 of Anopheles gambiae was found upregulated if infected with Plasmodium and in blood feeding [2,13]. As mentioned in the introduction PRS1 also relocated to vesicles in salivary glands and midgut in infection. Likewise, FgDM9-1 when tested in insect cells reacted by relocalization to vesicles following cellular stress in form of heat, exposure to bacteria (Pseudomonas aeruginosa and Escherichia coli) or incubation with chloroquine [1]. In Crassostrea gigas CGL1 was abundant in hepatopancreas, mantle and hemocytes. In hemocytes challenged with bacteria relocalization of CGL1 from the cell membrane to the cytoplasm was observed [6]. The authors speculated that proteins carrying DM9 domains provide an important defense mechanism against microbe invasion. This function is questionable for FgDM9-1 for 2 reasons; firstly Fasciola gigantica should not encounter bacteria in host liver and bile ducts; secondly the protein was detected in the parenchyma, an internal tissue without direct host contact. The experimental observations might overstate a protective function when the basic function might be in general cellular transport of molecules with mannose residues. Further experiments would be necessary to identify these molecules in Fasciola and other trematodes like Opisthorchis.

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CONFLICT OF INTEREST

We declare that we have no conflict of interest related to this work.

REFERENCES

1. Phadungsil W, Smooker PM, Vichasi-Grams S, Grams R. Characterization of a Fasciola gigantica protein carrying two DM9 domains reveals cellular relocalization property. Mol Biochem Parasitol 2016; 205: 6-15. https://doi.org/10.1016/j.molbiopara.2016.02.008
2. Chentemps T, Mitri C, Perrot S, Sautereau J, Jacques IC, Thierry I, Bourguin C, Rosinski-Chupin I. Anopheles gambiae PRS1 modulates Plasmodium development at both midgut and salivary gland steps. PLoS One 2010; 5: e11538. https://doi.org/10.1371/journal.pone.0011538
3. Labbunnuang N, Phadungsil W, Tesana S, Smooker PM, Grams R. Similarity of a 16.5kDa tegumental protein of the human liver fluke Opisthorchis viverrini to nematode cytoplasmic motility protein. Mol Biochem Parasitol 2016; 207: 1-9. https://doi.org/10.1016/j.molbiopara.2016.04.002
4. Grant RP, Buttery SM, Ekman GC, Roberts TM, Stewart M. Structure of MFP2 and its function in enhancing MSP polymerization in Ascaris sperm amoeboid motility. J Mol Biol 2005; 347: 583-595. https://doi.org/10.1016/j.jmb.2005.01.054
5. Utuno H, Matsuyma K, Tsuji Y, Goda S, Hiemori K, Tateno H, Hirabayashi J, Hatakeyama T. Identification, characterization, and X-ray crystallographic analysis of a novel type of mammose-specific lectin CGL1 from the pacific oyster Crassostrea gigas. Sci Rep 2016; 6: 29135. https://doi.org/10.1038/srep29135
6. Jiang S, Wang L, Huang M, Jia Z, Weinert T, Warkentin E, Liu C, Song X, Zhang H, Witt J, Qiu L, Peng G, Song L. DM9 domain containing protein functions as a pattern recognition receptor with broad microbial recognition spectrum. Front Immunol 2017; 8: 1607. https://doi.org/10.3389/fimmu.2017.01607
7. Sumner JB, Howell SE. Identification of Hemagglutinin of Jack Bean with Concanavalin A. J Bacteriol 1936; 32: 227-237. https://doi.org/10.1128/JB.32.2.227-237.1936
8. Goldstein II, Hollerman CE, Smith EE. Protein-carbohydrate interaction. II. Inhibition studies on the interaction of concanavalin A with polysaccharides. Biochemistry 1965; 4: 876-883. https://doi.org/10.1021/bi00881a013
9. Derewenda Z, Yariv J, Hellmwell JR, Kalb AJ, Dodson EJ, Papiz MZ, Wan T, Campbell J. The structure of the saccharide-binding site of concanavalin A. EMBO J 1989; 8: 2189-2193. https://doi.org/10.1002/jemt.22407
10. Waterhouse A, Bertoni M, Bienert S, Studer G, Tauriello G, Gumienny R, Heer FT, de Beer TAP, Rempfer C, Bordoli L, Lepore R, Schwede T. SWISS-MODEL: homology modelling of protein structures and complexes. Nucleic Acids Res 2018; 46: 296-303. https://doi.org/10.1093/nar/gky427
11. Rice P, Longden I, Bleasby A. EMBOSS: the European Molecular Biology Open Software Suite. Trends Genet 2000; 16: 276-277. https://doi.org/10.1016/S0168-9525(00)02042-2
12. Adamović I, Malinovská I, Wimmerová M. New sensitive detection method for lectin hemagglutination using microscopy. Micrsc Res Tech 2014; 77: 841-849. https://doi.org/10.1002/jemt.22407
13. Rosinski-Chupin I, Briolay J, Brouilly P, Perrot S, Gomez SM, Chertemps T, Roth CW, Keime C, Gandrillon O, Couble P, Brey PT. SAGE analysis of mosquito salivary gland transcriptomes during *Plasmodium* invasion. Cell Microbiol 2007; 9: 708-724. https://doi.org/10.1111/j.1462-5822.2006.00822.x

14. Beitz E. TEXshade: shading and labeling of multiple sequence alignments using LATEX2 epsilon. Bioinformatics 2000; 16: 135-139. https://doi.org/10.1093/bioinformatics/16.2.135

15. Pettersen EF, Goddard TD, Huang CC, Meng EC, Couch GS, Croll TL, Morris JH, Ferrin TE. UCSF ChimeraX: Structure visualization for researchers, educators, and developers. Protein Sci 2021; 30: 70-82. https://doi.org/10.1002/pro.3943