

This is the peer-reviewed version of the following article:

Raschdorf O., Bonn F., Zeytuni N., Zarivach R., Becher D., and Schüler D. (2018). A quantitative assessment of the membrane-integral sub-proteome of a bacterial magnetic organelle. Journal of Proteomics 172:89-99,

which has been published in final form at https://doi.org/10.1016/j.jprot.2017.10.007



This work is licensed under

<u>Creative commons Attribution-Noncommercial-NoDerivatives 4.0 International License</u>

A quantitative assessment of the membrane-integral sub-proteome

2	of a bacterial magnetic organelle
3	
4	Oliver Raschdorf ^{1a} , Florian Bonn ^{2b} , Natalie Zeytuni ^{3c} , Raz Zarivach ³ , Dörte Becher ²
5	and Dirk Schüler* ^{1,4}
6	Affiliations:
7	1: Department of Microbiology, Ludwig Maximilian University of Munich, Germany
8	2: Department of Microbiology, Ernst Moritz Arndt University of Greifswald,
9	Germany
10	3: Department of Life Sciences, The National Institute for Biotechnology in the
11	Negev and Ilse Katz Institute for Nanoscale Science and Technology, Ben-Gurion
12	University of the Negev, Beer Sheva, Israel
13	4: Department of Microbiology, University of Bayreuth, Germany
14	
15	* Corresponding author
16	^a Current address: Thermo Fisher Scientific (formerly FEI Company), Eindhoven,
17	Netherlands
18	^b Current address: Institute of Biochemistry II, Goethe University School of
19	Medicine, Frankfurt am Main, Germany.
20	^c Current address: Department of Biochemistry and Molecular Biology, University of

21

British Columbia, Canada

Keywords

22

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

23 Bacterial organelles; Magnetosomes; Membrane integral sub-proteome; Protein

24 quantification

Abstract

Magnetotactic bacteria produce chains of complex membrane-bound organelles that direct the biomineralization of magnetic nanoparticles and serve for magnetic field navigation. These magnetosome compartments have recently emerged as a model for studying the subcellular organization of prokaryotic organelles. Previous studies indicated the presence of specific proteins with various functions in magnetosome biosynthesis. However, the exact composition and stoichiometry of the magnetosome subproteome have remained unknown. In order to quantify and unambiguously identify all proteins specifically targeted to the magnetosome membrane of the Alphaproteobacterium Magnetospirillum gryphiswaldense, we analyzed the protein composition of several cellular fractions by semi-quantitative mass spectrometry. We found that nearly all genuine magnetosome membrane-integral proteins belong to a well-defined set of previously identified proteins encoded by gene clusters within a genomic island, indicating a highly controlled protein composition. Magnetosome proteins were present in different quantities with up to 120 copies per particle as estimated by correlating our results with available quantitative Western blot data. This high abundance suggests an unusually crowded protein composition of the membrane and a tight packing with transmembrane domains of integral proteins. Our findings will help to further define the structure of the organelle and contribute to the elucidation of magnetosome biogenesis.

Significance

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

Magnetosomes are one of the most complex bacterial organelles and consist of membrane-bounded crystals of magnetic minerals. The exact composition and stoichiometry of the associated membrane integral proteins are of major interest for a deeper understanding of prokaryotic organelle assembly; however, previous proteomic studies failed to reveal meaningful estimations due to the lack of precise and quantitative data, and the inherently high degree of accumulated protein contaminants in purified magnetosomes. Using a highly sensitive mass spectrometer, we acquired proteomic data from several cellular fractions of a magnetosome producing magnetotactic bacterium and developed a comparative algorithm to identify all genuine magnetosome membrane-integral proteins and to discriminate them from contaminants. Furthermore, by combining our data with previously published quantitative Western blot data, we were able to model the protein copy number and density within the magnetosome membrane. Our results suggest that the magnetosome membrane is specifically associated with a small subset of integral proteins that are tightly packed within the lipid layer. Our study provides by far the most comprehensive estimation of magnetosomal protein composition and stoichiometry and will help to elucidate the complex process of magnetosome biogenesis.

Highlights

68

71

72

73

74

75

- First quantitative and unbiased assessment of the membrane-integral sub-70 proteome of bacterial organelles (magnetosomes).
 - A comparative algorithm allows to distinguish genuine magnetosome integral protein from contaminants acquired during the isolation process, and to identify putative novel magnetosome membrane constituents.
 - Surface modelling suggests that a very specific subset of integral proteins is highly enriched and tightly packed within the magnetosome membrane.

Introduction

77

78 The Alphaproteobacterium Magnetospirillum gryphiswaldense and related 79 magnetotactic bacteria form intracellular, membrane-bounded crystals of a magnetic 80 mineral, the magnetosomes, which serve as magnetic sensors to help to direct 81 bacterial swimming towards growth-favoring suboxic zones in the sediments of 82 natural waters [1]. Magnetosome biosynthesis comprises the invagination of 83 magnetosome membrane vesicles from the cytoplasmic membrane [2,3], in which 84 conditions are properly controlled for the biomineralization of nano-sized crystals of 85 the iron oxide magnetite. Nascent magnetosomes are then aligned into linear chains 86 along cytoskeletal filaments to achieve one of the highest structural levels found in 87 prokaryotic cells [4]. 88 Magnetosome biosynthesis is thought to involve the sorting of a complex set of 89 proteins to the magnetosome membrane [3–5]. First comparative analyses suggested 90 the presence of specific magnetosome membrane proteins in various quantities which 91 co-purified with magnetosome particles isolated by magnetic collection and 92 ultracentrifugation [6–8]. Attempts to assess the magnetosome membrane proteome 93 of *M. gryphiswaldense* by denaturing one- and two-dimensional gel electrophoresis 94 followed by Edmann degradation and mass spectroscopy in combination with 95 comparative genomic analysis identified a set of about 25 bona fide magnetosome 96 proteins termed Mam (magnetosome membrane) and Mms (magnetosome membrane 97 specific) [7,9,10] which are thought to have key functions in magnetosome 98 membrane biogenesis, iron transport, nucleation and crystallization of magnetite as 99 well as the assembly of magnetosome chains [4]. The corresponding genes are 100 clustered within four operons of a hypervariable genomic magnetosome island (MAI), namely mamABop (17 genes), mamGFDCop (4 genes), mms6op (4 genes) 101

and mamXYop (4 genes) [10,11]. Other genes within the MAI but located outside these clusters were later also implicated in magnetosome formation (mamF2, mamD2, feoAB1, mamW; [10], [R. Uebe, manuscript in preparation]. However, in addition to bona fide magnetosome proteins, these MAI gene clusters also predict a number of further proteins which remained undetected in previous approaches, and it is unknown whether those represent genuine integral magnetosome membrane constituents or are just loosely attached, whether they are targeted exclusively to this compartment or also present in the cytoplasmic membrane, or in some cases, whether they are expressed at all. In addition, a multitude of other proteins encoded outside the well-established MAI gene clusters were found to co-purify with isolated magnetosomes [8,9,12], and so far it has yet not been resolved if some of these proteins represent either further functional integral constituents native to the magnetosome membrane, or contaminants that become bound upon cell disruption and isolation [8,13]. Moreover, previous proteomic approaches indicated that Mam and Mms proteins are present in the magnetosome membrane in vastly different quantities. However, attempts to estimate the abundance of several integral magnetosome membrane proteins by Coomassie-stained band intensities in protein gels [9] were inherently inaccurate. Thus, the complete protein complement of the magnetosome membrane still remains unknown, and it is not understood how and in which stoichiometry magnetosome proteins are assembled to form the structural framework required for biomineralization and organization of functional magnetic organelles. In this study we performed a highly sensitive, semi-quantitative mass spectrometry analysis of purified magnetosomes and several other cellular fractions and used a comparative algorithm to reveal the genuine membrane-integral magnetosome sub-

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

proteome and to confidently estimate the relative abundances of the individual
proteins.
With one exception we detected all previously predicted Mam and Mms proteins and
demonstrate that several of them are highly and specifically enriched in the
magnetosome membrane. Furthermore, we identified several novel putative genuine
magnetosome-membrane proteins, of which one (MGR_4114) is encoded within the
MAI. Our results also indicate that most of the proteins detected within the
magnetosome-membrane fraction, but encoded outside the MAI are likely
contaminants from other cellular compartments.
Correlation of our semi-quantitative proteomic data with available quantitative
Western blot data allowed us to approximate the absolute copy numbers of
magnetosome membrane proteins within the organelle. The magnetite-nucleating
Mms6 was identified as the most abundant membrane-integral magnetosome protein,
followed by MamC and MamD, together accounting for >40% of all genuine
magnetosome membrane proteins. Using the predicted topology of magnetosome
proteins, we could further estimate the membrane coverage of integral proteins,
which hints towards an unusually crowded protein organization within the
magnetosome membrane. In summary, our data allowed the most accurate estimation
of protein composition of the complex magnetosome membrane up to date and will
contribute to uncover the processes involved in biogenesis of this sophisticated
bacterial organelle.

Experimental Procedures

149	Cultivation and cell harvesting of M. gryphiswaldense, C_{mag} determinations
150	Bacterial strains and plasmids used in this study are listed in suppl. Table S 7. E. coli
151	strains were cultivated in lysogeny broth (LB) medium. When necessary, kanamycin
152	(Km) was added to 25 μg mL ⁻¹ . E. coli BW29427 and WM3064 cultures were
153	supplemented with 1 mM DL- α , ϵ -diaminopimelic acid. Media were solidified by
154	addition of 1.5% (w/v) agar. M. gryphiswaldense cultures were grown at 30°C in
155	modified flask standard medium (FSM) [14]. When appropriate, Km was added to 5
156	μg mL ⁻¹ . Optical density (OD) and magnetic response (C _{mag}) of exponentially
157	growing cultures were measured photometrically at 565 nm as described previously
158	[15]. Conjugations of plasmids were performed essentially as described earlier
159	[16,17].
160	Cellular fractionation and purification
161	Cultivation and all further fractionation steps were performed in independent
162	triplicates as described in the workflow chart of Figure 1. M. gryphiswaldense was
163	cultivated and scaled up to 4,5 L culture in closed 5 L-Schott bottles (500 ml air in
164	headspace) over-night at 30°C and harvested by centrifugation (10,000 g, 15 min,
165	4°C). The cell pellets were washed in buffer containing 20 mM HEPES (pH 7.4) and
166	5 mM EDTA, and frozen at -20°C. All further steps were carried out at 4°C. The cell
167	pellets obtained from cell harvesting were resuspended in buffer containing 50 mM
168	HEPES (pH 7.4), 1 mM EDTA and complete protease inhibitor (Roche, Germany)
169	and lysed by a high-pressure cell disruption system. Cellular storage
170	polyhydroxybutyrate (PHB) granules were removed by centrifugation (210 g, 10 min
171	4°C) of the lysate. The lysate was applied on MACS cell separation column type CS
172	(Miltenyi Biotec, Germany), magnetized with two neodymium-iron-boron cube

magnets (gravity flow). The flow-through was applied a second time on the column
and then collected as total nonmagnetic lysate (non-mag). The fraction bound to
the separation column was washed with 50 mL extraction buffer [10 mM HEPES
(pH 7.4), 1 mM EDTA, 0.1 mM PMSF], two times 50 mL salt buffer [10 mM
HEPES (pH 7.4), 1 mM EDTA, 200 mM NaCl, 0.1 mM PMSF] and again 50 mL
extraction buffer by gravity flow. The magnets were removed; the magnetic fraction
eluted from the column with approx. 10 mL H_2O and supplemented to final HEPES
(pH 7.4), EDTA and PMSF concentrations of the extraction buffer. An
ultracentrifugation tube was filled with 60% (w/w) sucrose (in extraction buffer) and
overlaid with the magnetic fraction. After ultracentrifugation (100,000 g, 1.5 h),
pellet was resuspended in 2 mL extraction buffer as magnetically separated
magnetosome fraction (mag).
The total nonmagnetic cellular lysate was centrifuged for 1 h at 100,000 g and the
membrane pellet resuspended (central small white PHB pellet was omitted) and
incubated in carbonate buffer (200 mM Na ₂ CO ₃ , 10 mM EDTA, 1 mM PMSF, pH
11.0) for 30 min under mild shaking. After centrifugation for 1 h at 100,000 g, the
pellet was resuspended in high-salt buffer (20 mM Tris, 1 M NaCl, 10 mM EDTA, 1
mM PMSF, pH 7.5) and incubated under mild shaking for 30 min. After
centrifugation for 1 h at 100,000 g, the pellet was resuspended in 50 mM TEAB (pH
7.8) and immediately pelleted for 1 h at 100,000 g. The pellet formed the total
nonmagnetic membrane fraction (mem).
The magnetically separated magnetosomes (mag) were centrifuged for 30 min at
100,000 g and the pellet resuspended in carbonate buffer. Subsequent purification
was performed as described for the membrane fraction, with 30 min centrifugation
runs between washes. The resulting pellet formed the stringently washed
magnetosome (mag.str) fraction (Figure 1).

SDS-PAGE, tryptic digestion and mass spectroscopy analysis
All triplicate fractions were treated independently. For sodium dodecyl sulfate
polyacrylamide gel electrophoresis (SDS-PAGE), all liquid samples were
supplemented and all pelleted samples were dissolved in 2x SDS sample buffer
[0.125 M Tris (pH 6.8), 4% SDS, 2% glycerol, 10% 2-mercaptoethanol, 0.004%
(w/v) Bromophenol blue] and heated at 60°C for 5 min. Appropriate amounts of
samples were determined empirically (Fig S 2). Electrophoresis of the protein
samples was performed on 12% polyacrylamide gels. Staining, in gel tryptic
digestion and LC-MS/MS was performed according to [18] with minor
modifications. In brief, the in-gel digested peptides were separated with an easy nLC
2 (Thermo Fisher Scientific, MA, USA) column and analyzed with an LTQ Orbitrap
Velos (Thermo Fisher Scientific). The 20 most intense precursor ions of each full
scan were selected for collision induced dissociation fragmentation. A list of all open
reading frames (ORFs) from the draft genome sequence of M. gryphiswaldense [10],
was used as target database, supplemented with entries of recently assigned ORFs.
The resulting output files were compiled with Scaffold 4 (Proteome Software, OR,
USA). Proteins were only considered as identified if at least two unique peptides,
matching solid quality criteria ($\Delta cN > 0.1$ and $XCorr > 2.2; 3.3; 3.7$ for doubly,
triply, or higher charged peptides) have been assigned, resulting in a false positive
rate below 0.2% on protein level. Only two reverse decoy peptides were assigned in
the screen, indicating good filter criteria. Spectral counts for these two peptides were
omitted from further analysis. Spectral counts from known contaminants (e.g. human
source, trypsin) were also excluded from further analysis

222	Proteinase K membrane shaving
223	If not otherwise noted, all steps were carried out at 4°C. A schematic description of
224	the process is presented in Figure 3. A 1 mL batch of magnetically separated
225	magnetosomes (mag) was centrifuged for 30 min at 100,000 g. The pellet was
226	resuspended in carbonate buffer and incubated for 1 h under mild shaking. Urea was
227	added to final concentration of 8 M. For protein reduction, tris(2-
228	carboxyethyl)phosphine hydrochloride was added to final concentration of 5 mM and
229	sample incubated for 45 min at 60°C. Alkylation was performed by addition of 10
230	mM iodoacetamide and incubation for 15 min at room temperature in the dark.
231	Proteinase K was added to a final concentration of 3 $\mu g/ml$ and the sample was
232	proteolytically digested for 15 h at 37°C under mild shaking. Samples were
233	supplemented with 5% acetonitril, cooled down on ice and centrifuged for 1 h at
234	100,000 g. Supernatant was removed and pellet overlaid with 50 mM TEAB solution
235	before centrifugation for 1 h at 100,000 g. The resulting pellet was frozen at -70°C.
236	Pellet was resuspended in 180 μL of digestion buffer [50 mM TEAB, pH 7.8, 0.5%
237	RapiGest (Waters, MA, USA)] and incubated for 30 min at 30°C under mild shaking.
238	$6~\mu g$ of Chymotrypsin and $10~mM~CaCl_2$ were added and solubilized sample digested
239	for 7 h at 30°C with mild shaking. HCl was added to final concentration of 250 mM
240	and sample incubated at 37°C for 45 min to precipitate detergent. Sample was
241	repetitively centrifuged for 12 min at 20,000 g, until no magnetosome and membrane
242	pellet was visible. The supernatant formed the shaved magnetosome
243	transmembrane peptide fraction and was analyzed by LC-MS/MS as described in
244	[19].

245 Mass spectrometry proteomics Raw data deposition

The mass spectrometry proteomics data have been deposited to the

247 ProteomeXchange Consortium via the PRIDE [20] partner repository with the

248 dataset identifier PXD006166.

249 Data analysis

246

253

254

255

256

257

258

250 All calculations described in the main text were performed in Microsoft Excel and

are included in the supplements (suppl. File 1). To develop our working model, we

252 made the following assumptions:

(1) The **relative abundance** of a single protein in a complex sample can be estimated by normalizing the assigned peptide spectra for this specific protein with the total number of peptide spectra measured in this sample and with the molecular weight (MW) of the protein. This normalization allows a size-independent comparison of protein abundance over several fractions. The estimated abundance (A) of protein N is defined by

$$A_N = \frac{SpC_N}{\left[\left(\sum_{i=1}^n SpC_i\right) \times MW_N\right]}$$

whereas

N is the protein index

SpC is the number of unique peptide spectra matching the protein

262 MW is the molecular weight of the protein

n is the total number of proteins identified in the screen of a single sample

For further calculations, the A_N of all detected proteins was independently

determined in all samples and averaged over the three replicate fractions of a sample

266 type.

- (2) A genuine integral magnetosome membrane protein has to fulfill the following conditions:
 - a. The protein is relatively more enriched in the membrane fraction than in the total non-magnetic lysate, *i.e.* it is a membrane protein.
- 271 Determined by:

268

269

270

- 272 $A_{\text{mem(N)}}/A_{\text{non-mag(N)}} \ge 1$
- b. The protein is comparatively more [or by the factor of 'f' much more] 273 274 enriched in the magnetosome membrane than in the non-magnetic 275 membrane fraction of the cell. Determined by:
- 276 $A_{\text{mag(N)}}/A_{\text{mem(N)}} > 1 > f$
- 277 c. The protein becomes more highly enriched in magnetosomes that 278 were depleted from associated proteins and contaminations by 279 stringent washing. Determined by:
- 280 $A_{\text{mag.str}(N)}/A_{\text{mag}(N)} \ge 1$
- 281 d. Optional condition: The protein has comparatively very high relative 282 abundance in the magnetosome membrane fraction. Determined by: $A_{\text{mag.str(N)}}/A_{\text{mag.str(MamC)}} > T$ (The estimated abundance of the protein 283 284 has to at least meet threshold T, when compared to a known highly 285 abundant magnetosome protein, here MamC)
- 286 Calculations for magnetosome protein abundance and surface coverage model 287 Based on quantitative Western blots with MamC-GFP labeled magnetosomes and the 288 correlation with magnetite crystal size and density, a conservatively estimated 289 number of approximately 100 molecules (estimated range: 80 - 250) of MamC were 290 suggested for an average-sized magnetosome [21]. By calculating the A_{mag,str(N)}/A_{mag,str(MamC)} value for every protein of interest, we estimated its copy

number in an average magnetosome, assuming a MamC reference copy number of 100. To calculate the transmembrane domain (TMD) coverage of the magnetosome membrane, we made the following assumptions: i) A simple transmembrane helix (TMH), perpendicularly inserted into the lipid bilayer, has a cross-section diameter of at least 1.1 nm [22]. This is a very conservative assumption, since contributing large amino acid residues and different insertion angles might increase the crosssection diameter ii) TMHs are surrounded by boundary lipids that interact with the hydrophobic protein domain. The major phospholipids in the cytoplasmic and magnetosome membrane of *M. gryphiswaldense* are phosphatidylethanolamine (PE) and phosphatidylglycerol (PG) [9]. The lipid head cross-section area and the diameter of dilauroyl-PE and dilauroyl-PG are 0.39 nm² (0.70 nm) and 0.43 nm² (0.74 nm), respectively [23]. On average, the diameter of a lipid head group in the magnetosome membrane can therefore be estimated to be 0.72 nm. Hence, an annular boundary lipid shell would increase the diameter of an embedded TMH to 2.5 and 4.0 nm for one and two rings of boundary lipids, respectively. iii) TMHs of multi-membrane spanning proteins are more packed and without internal lipid boundary layers. According to Jacobson et al. [22], a diameter of 2.4 nm can be assumed for the whole TMD of a tetraspan-protein, and a diameter of 3.2 nm for a heptaspan-protein, excluding boundary lipids (Figure 3). We interpolated these values for magnetosome membrane proteins that exhibit 1 to 18 predicted TMH [5] by power regression. The average diameter of a magnetosome vesicle from M. gryphiswaldense is 45.5 nm [3]; the spherical surface area therefore can be calculated to be approx. 6450 nm². Taking into account the number of predicted TMHs and the here estimated absolute copy numbers of genuine magnetosome proteins, we calculated the integral protein occupancy of the membrane (surface) for both the

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

312

313

314

315

317	scenarios that all TMH are isolated and that TMHs of a single protein form a packed
318	TMD (Figure 3).
319	To control our quantification results, we also used the protein abundance index (PAI)
320	as an alternative method for quantifications [24]. The method does not take into
321	account the molecular weight of the protein for normalization, but rather the number
322	of peptides that are theoretically generated by the utilized protease (here: trypsin) and
323	also excludes peptides that are too small or big to be measured by mass spectroscopy.
324	We calculated three different PAI values for all assigned Mam and Mms proteins
325	(PAI I: assuming theoretical tryptic peptides between 600 and 5000 Da, PAI II:
326	assuming theoretical tryptic peptides with 7 to 25 amino acids and maximum
327	molecular weight of 5000 Da and PAI III: assuming the number of tryptic peptides
328	that were actually detected in our analysis). See also suppl. File 1.
329	Molecular and genetic techniques
330	Oligonucleotides were purchased from Sigma-Aldrich (Germany) and are listed in
331	suppl. Table S 8. Plasmids were constructed by standard recombinant techniques
332	using enzymes from Thermo Fisher Scientific and Agilent Technologies (CA, USA)
333	and confirmed using BigDye terminator v3.1 chemistry on an ABI 3700 capillary
334	sequencer (Thermo Fisher Scientific). All plasmids used in this study are listed in
335	suppl. Table S 7.
336	Plasmids pOR129 and pOR156 for markerless in-frame deletion of MGR_3691 and
337	the MGR_4114 operon respectively, were created by PCR amplification and fusion
338	of approximately 750 bp regions up- and downstream of the target sequences and
339	cloning into pORFM. Genes were deleted as described in [25].
340	Magnetosome proteins were C- or N-terminally fused to enhanced green fluorescent
341	protein (EGFP) and (over)expressed under control of the strong $P_{mamDC45}$ [21]

promotor either from replicative plasmids or by transposon mediated random integration in the genome. In all cases, the two fusion proteins were separated by a 25 amino acid alpha-helical linker region [LA(EAAAK)₄AAA] (HL) [3,26]. Alternatively, in-frame genomic fusions were constructed. Replicative plasmids pOR079, pOR089 and pOR099 were constructed by replacing mamI in pOR075 by PCR-amplified mamW, mamR or mms6 sequences, respectively, using restriction digestion. Similarly, pOR085, pOR087, pOR088, were constructed by replacing mamQ in pOR086 by mmsF, mamR and mamE sequences, respectively. The P_{mamDC} mamE-HL-egfp sequence form pOR088 was further cloned into the transposable plasmid pBAM-1 by restriction digestion, creating pOR148. A chromosomal mamA-GFP fusion was generated as described in [25], using plasmid pOR068. The plasmid was created by exchanging flanking regions of mamC in pFM236, by approximately 750 bp homologous flanking regions of *mamA* using restriction digestion. Microscopy methods For fluorescence microscopy, 3 µl samples of M. gryphiswaldense over-night cultures were immobilized on 1% (w/v) agarose pads with FSM medium salts. The samples were imaged with an BX81 microscope (Olympus, Japan) equipped with a 100×UPLSAPO100XO 1.4NA objective and an Orca-ER camera (Hamamatsu, Japan) and appropriate filer sets using Olympus Xcellence software. For transmission electron microscopy (TEM), unstained formaldehyde-fixed (0.075% w/v) M. gryphiswaldense cells were absorbed on carbon coated copper grids. Bright field TEM was performed on a Phillips (Netherlands) CM200 instrument using an accelerating voltage of 160 kV. Images were captured with an Eagle 4k CCD camera using EMMenu 4.0 (Tietz, Germany).

366

342

343

344

345

346

347

348

349

350

351

352

353

354

355

356

357

358

359

360

361

362

363

364

Results and Discussion

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

382

383

384

385

386

387

388

389

390

391

392

393

The four following different cellular fractions from M. gryphiswaldense were prepared for comparative mass spectroscopy analysis: (1) Magnetically separated magnetosomes (mag) following previously suggested protocols [27]; (2) stringently washed magnetosomes (mag.str), additionally washed in high-salt and alkaline carbonate buffer to deplete contaminating proteins; (3) total **non-magnetic lysate** (non-mag) obtained as flow-through of magnetic column separation; (4) total enriched non-magnetic membrane fraction (mem), washed in high-salt and alkaline carbonate buffer (Figure 1). Analysis of the mass spectroscopy data in total showed peptides of 2237 unique proteins in all fractions and replicates, which would account for approximately 53% off all predicted open reading frames (ORFs) of M. gryphiswaldense [28], and over 1000 proteins in the magnetosome fraction, indicating that the number of magnetosome-associated proteins would be unreasonably high without adequate filter algorithms to identify the most realistic subset of genuine magnetosome-associated proteins. Peptides of 1135 proteins were detected in magnetically separated magnetosomes, 1027 proteins in the stringently washed magnetosome fraction, 2031 proteins in the total non-magnetic fraction and 1305 proteins in the enriched non-magnetic membrane fraction.

Establishing and evaluating criteria to identify the genuine integral magnetosome

membrane sub proteome

Genuine integral magnetosome proteins are bound to the magnetosome membrane in vivo and are specifically and exclusively enriched in this compartment. We developed a working model to identify these genuine integral magnetosome proteins and to discriminate them from contaminates by simply comparing and weighting the estimated relative abundance A of all proteins found within four cellular fractions. To fulfill our criteria, the protein has to be (I) a membrane-associated protein (determined by $A_{mem}/A_{non-mag}$ -ratio of ≥ 1), (II) strongly enriched in the magnetosome

- membrane compared to the non-magnetic membrane fraction (determined by
- $A_{\text{mag}}/A_{\text{mem}}$ -ratio of > 1), and (III) would not become depleted from the magnetosome
- 396 by stringent washing (determined by $A_{mag.str}/A_{mag}$ -ratio of ≥ 1) (For more
- information, see data analysis section in experimental procedures).
- In total, only 81 proteins (of which 23 were Mam and Mms annotated) were assigned
- 399 genuine integral magnetosome membrane proteins if the parameters of our model
- $400 \quad \text{were set to} \quad A_{mem}/A_{non\text{-}mag} \geq 1, \quad A_{mag}/A_{mem} > 1 \quad \text{and} \quad A_{mag,\text{str}}/A_{mag} \geq 1; \quad (Table \quad 1).$
- 401 Application of each individual criteria contributed to the exclusion of proteins from
- 402 the list of genuine magnetosome proteins, i.e. without taking into account every term,
- 403 the number of assigned magnetosome proteins, especially those that are not encoded
- 404 by the mam and mms operons, would have been unreasonably high (up to 651)
- 405 (Table 1 and
- 406 suppl. Table S 1). On the other hand, when $A_{mem}/A_{non-mag}$ and $A_{mag.str}/A_{mag}$ were set
- above the value of ≥ 1 , e.g. to ≥ 3 and ≥ 2 , respectively, the number of assigned
- 408 proteins, and especially of those encoded outside the well-established mam and mms
- 409 gene clusters shrank dramatically, indicating that the two terms should not
- 410 exceed ≥ 1 (
- 411 suppl. Table **S 1**).
- 412 Accordingly, the total number of assigned proteins further decreased when the
- 413 A_{mag}/A_{mem} ratio was set more stringently, while the number of assigned Mam/Mms
- 414 magnetosome membrane proteins only decreased by two when A_{mag}/A_{mem} was
- changed from > 1 to > 4 (Table 1 and suppl. Table S 2), indicating that the more
- 416 stringent conditions are still sufficient to identify experimentally confirmed
- 417 magnetosome membrane proteins. Therefore, parameter sets of $A_{\text{mem}}/A_{\text{non-mag}} \ge 1$,
- 418 $A_{mag}/A_{mem} \ge 4$ and $A_{mag.str}/A_{mag} \ge 1$ seemed to be appropriate to predict the most
- 419 likely genuine magnetosome proteins.
- 420 By comparing the estimated abundance of individual proteins within the stringently
- washed magnetosome fraction, the set of genuine magnetosome proteins might be

422 further refined. MamC was previously suggested to be the most abundant protein in 423 the magnetosome membrane [7,29]. An A_{mag,str(N)}/A_{mag,str(MamC)}-threshold of 0.01 would indicate that for 100 estimated copies of MamC, at least one copy of the 424 425 protein of interest N has to be present in the magnetosome membrane. The effect of 426 supplementing the aforementioned conditions by the filter $A_{mag.str(N)}/A_{mag.str(MamC)} > T$ 427 (T of 0.1, 0.01 or 0.001) to eliminate low abundant proteins is visualized in suppl. 428 Table S 3. A threshold T of 0.1 further reduced the number of predicted non-429 magnetosome membrane proteins by 38 for $A_{mem}/A_{non-mag} \ge 1$, $A_{mag}/A_{mem} \ge 1$, 430 $A_{\text{mag.str}}/A_{\text{mag}} \ge 1$ and by 4 for the more restrictive $A_{\text{mem}}/A_{\text{non-mag}} \ge 1$, $A_{\text{mag}}/A_{\text{mem}} \ge 4$, 431 $A_{\text{mag.str}}/A_{\text{mag}} \ge 1$, while being alone not sufficient for adequate filtering (suppl. Table 432 S 3). In combination with the parameters $A_{mem}/A_{non-mag} \ge 0$, $A_{mag}/A_{mem} \ge 4$, $A_{mag.str}/A_{mag} \ge 1$, a $A_{mag.str}/A_{mag.str(MamC)}$ threshold T of 0.01 predicted only 30 genuine 433 434 magnetosome membrane proteins, of which 22 were previously identified Mam/Mms 435 proteins. These values are comparable to the effects of $A_{mem}/A_{non-mag} \ge 1$, 436 $A_{mag}/A_{mem} \ge 4$, $A_{mag.str}/A_{mag} \ge 1$ without MamC abundancy threshold and particular 437 interesting as an alternative because the $A_{mem}/A_{non-mag}$ coefficient could potentially 438 also exclude proteins that are so highly affine to the magnetosome membrane that 439 they are completely undetectable in the nonmagnetic membrane fraction. 440 Mam and Mms proteins comprise the major fraction of the genuine magnetosome 441 membrane proteome 442 With the exception of the small MamL, all other proteins encoded by the mam and 443 mms gene clusters were identified in our proteomic data, including the recently 444 identified MamF2 and MamD2 as well as MamX and MamI, which escaped 445 detection in previous proteomic studies [13]. MamL is a small (approx. 9 kDa) 446 protein that contains two predicted hydrophobic transmembrane domains. Although

447	two predicted tryptic peptides are in the theoretically detectable mass range, they
448	were not detected in any of the samples of this study. However, as indicated by
449	studies with MamL-GFP fusions and the strong magnetosome phenotype of gene
450	deletion [3,30], the MamL protein is expressed and at least partially targeted to the
451	magnetosome membrane, but was also never detected in previous proteomic
452	analyses. In contrast, small proteins of comparable low mass such like MamR
453	(approx. 8 kDa) and MamI (approx. 7 kDa) were confidently identified in this study.
454	MamI also comprises two predicted trans-membrane domains, but more predicted
455	tryptic peptides than MamL. The absence of MamL peptides from the data is an
456	indication that for unknown reasons some proteins might be underrepresented or
457	false-negatives in our screen.
458	With the stringent parameter setting of $A_{mem}/A_{non-mag} \ge 1$, $A_{mag}/A_{mem} \ge 4$,
459	$A_{mag.str}/A_{mag} \ge 1$, the 21 assigned and also predicted genuine integral magnetosome-
460	membrane proteins were: MamE, MamD, Mms6, MamO, MamM, MamC, MamB,
461	MamY, MamP, MamF2, MmsF, MamF, MamS, MamH, MamT, MamZ, MamI,
462	MamN, MamW, MamX and MamG. The two proteins MamQ and Mms48
463	(MGR_4070) met $A_{mem}/A_{non\text{-}mag} \geq 1$ and $A_{mag.str}/A_{mag} \geq 1,$ but only showed an
464	$A_{\text{mag}}/A_{\text{mem}}$ value of 2.0 or 1.6 respectively, therefore barely escaping our
465	classification as genuine magnetosome membrane proteins. It is thus possible that
466	both proteins are indeed genuine magnetosome-membrane proteins that are similarly
467	abundant in the cytoplasmic membrane; however, with an $A_{mag.str}/A_{mag.str(MamC)}$ value
468	of 0.01, the abundance of Mms48 in the magnetosome membrane fraction is very
469	low (i. e. 100 times less abundant than MamC). In the case of MamQ, this is
470	consistent with the observation by microscopy that MamQ-GFP within the cells was
471	mostly localized in the CM rather than the magnetosomes [3].
472	Several other Mam/Mms proteins also did not meet the criteria for genuine

magnetosome-membrane proteins. MamA, for example, failed by two criteria: Although its calculated A_{mag}/A_{mem} value of 51.2 was the second highest in the whole experiment, it did not fulfill the membrane protein threshold $(A_{mem}/A_{non-mag} = 0.6)$ and was depleted from the magnetosome membrane in the purification process $(A_{mag.str}/A_{mag}=0.3)$. Consistent with its lack of predicted transmembrane helices (TMH), it is well-established that MamA only associates to the magnetosome surface and is not an integral part of the MM membrane, but becomes readily depleted with alkaline treatment [31–33]. Our results therefore are in good agreement with previous findings, and provide a further validation for the effectiveness of the selected parameter set. MamJ was excluded for the same reasons, however shows less clear parameter values ($A_{mem}/A_{non-mag}$ and $A_{mag.str}/A_{mag} = 0.9$). Since it also does not contain predictable transmembrane domains, MamJ in vivo is most likely strongly magnetosome-membrane associated, but not integral. The actin-like MamK protein was excluded since it was more abundant in the non-magnetic membrane (mem) than the magnetosome membrane (mag) and the total non-magnetic lysate (non-mag). The latter indicates that polymerized, high molecular weight MamK is either pelleted with the membrane fraction, or is bound to the cytoplasmic membrane. Although MamD2 (like MamF2) was highly enriched in the magnetosome membrane, the protein did not meet the criteria since it became depleted from stringently washed magnetosomes ($A_{\text{mag.str}}/A_{\text{mag}} = 0.6$. The relevance of this finding remains unclear, but might indicate that the protein is not an integral magnetosome membrane protein, despite of its two predicted transmembrane domains. On the contrary, the 8 kDa MamR lacks any predicted transmembrane domain. However, it was virtually exclusively found in the magnetosome fraction, and could not be

473

474

475

476

477

478

479

480

481

482

483

484

485

486

487

488

489

490

491

492

493

494

495

496

depleted. With a A_{mag.str(MamR)}/A_{mag.str(MamC)} value of 0.2, it was additionally highly abundant and therefore seems to be tightly magnetosome associated. Since it was undetectable in the non-magnetic membrane (mem) and almost undetectable in the soluble protein fraction (non-mag), MamR is an ambiguous case but formally had to be excluded. Although MamD and MamR are thus most likely no integral components, the proteins seem tightly bound to purified particles. Similar to Mms48, Mms36 was excluded because of its low A_{mag}/A_{mem} value of 1.0. Additionally, both proteins show a very low predicted abundance in the magnetosomes. Hence, Mms48 and Mms36, which were found to have a non-essential role in magnetite formation [34], rather are localized in the cytoplasmic membrane in vivo. The mamXY operon encoded protein FtsZm by far failed all criteria and was almost non-detectable in magnetosomes. It is therefore most likely also active in another cellular compartment in vivo, presumably the cytoplasm. Finally, the predicted soluble MamU was found to be expressed, but also failed to pass any of the set criteria and therefore is most likely not magnetosome associated in vivo, which is in line with the absence of a discernible magnetosome phenotype upon deletion of mamU [34]. The MAI-encoded iron transporter homologue FeoAB1 [R. Uebe, manuscript in preparation] was recently implicated in magnetite formation [35]. While FeoA1 was not found in our screen, FeoB1 was present with the same abundance in magnetic and non-magnetic membrane fraction ($A_{mag}/A_{mem} = 1.0$, $A_{mem}/A_{non-mag} = 2.7$, $A_{mag.str}/A_{mag} = 1.2$), and therefore not found to be specifically enriched in the magnetosome membrane, but might still be a potential constituent of it. The predominant localizations of many Mam/Mms proteins in M. gryphiswaldense and other magnetotactic bacteria were already assessed by tagging and fluorescence microcopy analysis in previous studies [e.g 25,36–38]. In order to corroborate the findings of the proteome analysis, we investigated the intracellular localization of

498

499

500

501

502

503

504

505

506

507

508

509

510

511

512

513

514

515

516

517

518

519

520

521

522

some of those Mam/Mms proteins whose localization was not yet assessed previously in M. gryphiswaldense. GFP-labeled Mms6 and MmsF predominantly showed a strong linear fluorescent signal within the cells, resembling the organization of the magnetosome chain and indicating a strong enrichment in the magnetosome membrane (Figure 2A+B) as seen in our proteomic analysis. GFP-MamE also showed an accumulation of signal at mid-cell, however with higher cellular background (Figure 2C). Since MamW is only conserved in magnetospirilla, not part of one of the four major operons of the MAI, and its gene deletion did not shown any phenotype [13,30], its participation in magnetosome formation was yet not proved. The linear signal of the MamW-GFP fusion (Figure 2F) however is consistent with its proteomic detection and further suggests that MamW is specifically and genuinely magnetosome-integral. MamR and MamA GFP-fusions also showed a weak linear localization signal within the cell, along with a high cytoplasmic background (Figure 2D+E). These results corroborate the finding that both proteins were not assigned genuine integral magnetosome membrane proteins in this study, but were still found highly enriched in the magnetosome fraction, indicating strong magnetosome association. Given the high sensitivity of detection, it is highly likely that most of the 1000 proteins that were detected in the magnetosome fraction represent contaminations from other cellular compartments, resulting from unspecific binding during cell disruption and purification. Comparable to previous proteomic studies, predicted outer membrane proteins and ATPase subunits were some of the most abundant proteins in the magnetosome membrane fraction (mag) [7–9] (Table 2). However, these proteins are known to be among the most abundant cytoplasmic membrane proteins in all bacteria [39] and did not meet our stringent filter criteria for genuine magnetosome-membrane proteins, showing that some highly abundant proteins in the fraction contaminants. magnetosome are likely Next, we assessed if novel candidates identified in our screen for genuine

524

525

526

527

528

529

530

531

532

533

534

535

536

537

538

539

540

541

542

543

544

545

546

547

548

549

550

552 magnetosome-membrane proteins could be linked to magnetosome biosynthesis: 553 Besides the known Mam and Mms proteins discussed above, only few other proteins 554 met our filter criteria or were highly enriched in the magnetosome membrane (Table 555 1 and Fehler! Verweisquelle konnte nicht gefunden werden.), but are not 556 predicted to be magnetosome-associated (non Mam/Mms). Only one of those, the 557 small (7 kDa) hypothetical transmembrane protein MGR 4114, is conspicuously 558 encoded within the genomic magnetosome island, forming an operon together with 559 two additional hypothetical proteins and one protein with similarities to ParA/MinD-560 like ATPases. However, our deletion of the entire operon failed to cause a discernible 561 magnetosome phenotype (Fig S 3). Another candidate, MGR_3691 (also known as MM22 [9]) exhibited the 562 563 highest magnetosome enrichment of all proteins identified in our screen 564 $(A_{\text{mag}}/A_{\text{mem}} = 64.8)$, but was not assigned genuine integral magnetosome protein, as it 565 became depleted in stringently washed magnetosomes, and did not meet the integral 566 membrane protein threshold $(A_{\text{mag.str}}/A_{\text{mag}} = 0.5, A_{\text{mem}}/A_{\text{non-mag}} = 0.5)$. Indeed, 567 deletion of the gene did also not cause any magnetosome phenotype (568 Fig S 3), indicating that it has no important function in magnetosome formation. 569 Among the most abundant proteins, the previously identified Mms16 (MGR_0659) 570 was also identified, but not assigned genuine magnetosome protein (Table 2) due to 571 an A_{mem}/A_{non-mag} value of only 0.6 and an A_{mag}/A_{mem} value of 2.4. The protein was 572 previously implicated in magnetosome formation [40], but later in fact shown to 573 represent a phasin that rather functions in PHB metabolism of M. gryphiswaldense 574 [41]. This indicates that contaminations can be successfully uncovered by our 575 approach. While three of the other magnetosome assigned (according to our results), 576 but non-magnetosome predicted proteins showed a A_{mag.str(MamR)}/A_{mag.str(MamC)} below 577 0.01 (Table 2), five further proteins exhibited a value between 0.01 and 0.02. These 578 are the hypothetical proteins MGR_2833, MGR_0916 and MGR_2730 as well as the 579 histidine kinase MGR_0622 and the *ccb3*-type cytochrome oxidase maturation 580 protein MGR_2552 (Table 2). Since their abundance in the magnetosome membrane

581 is very low in comparison with MamC and currently no functional connection to 582 magnetosome formation is known for those proteins, it is highly likely that they 583 represent false positives with respect to identified genuine magnetosome-membrane 584 proteins. 585 Table 3 shows additional proteins that might be genuine magnetosome-membrane 586 proteins, identified using less stringent filter parameters ($A_{mem}/A_{non-mag} \ge 1$, 587 $A_{\text{mag}}/A_{\text{mem}} \ge 1$, $A_{\text{mag.str}}/A_{\text{mag}} \ge 1$), but with a predicted abundance of at least two 588 protein copies per magnetosome ($A_{\text{mag.str(N)}}/A_{\text{mag.str(MamC)}} \ge 0.02$). Besides MamQ, 589 these proteins comprise putative translocases, peptidases, hypothetical proteins, acyl-590 or glycosyl-transferases. However, only MamQ shows a reasonably high A_{mag}/A_{mem} 591 enrichment coefficient and also A_{mag.str}/A_{mag.str(MamC)} abundance. Similarly, ignoring 592 the A_{mag}/A_{mem} threshold, only MamR and a putative phasin could be additionally 593 associated with high A_{mag.str}/A_{mag.str(MamC)} abundance. 594 In summary, except few novel candidate proteins, of which only one showed higher 595 confidence (MGR_4114), it therefore appears that the specific integral 596 magnetosome-membrane sub-proteome is comprised of only previously predicted magnetosome-membrane proteins. 597 598 Proteinase K-shaved magnetosome membranes mostly contain MAI-encoded 599 Mam/Mms protein fragments 600 To obtain additional information about the integral magnetosome-membrane 601 proteome, we performed a Proteinase K membrane shaving assay on isolated 602 magnetosomes, which should cleave off and digest all external protein domains, 603 while buried transmembrane and membrane enclosed domains stay intact. Mass 604 spectroscopy analysis of shaved magnetosomes membranes assigned peptide spectra 605 to 61 proteins (suppl. Table S 4). Although we found previously undetected peptides

derived from predicted hydrophobic transmembrane helices of Mam and Mms proteins, most assigned spectra were from predicted soluble protein domains. However, among the 20 proteins with the highest exclusive un-normalized peptide spectra counts (SpC), 15 were Mam and Mms proteins (suppl. Table S 4). While 19% of all spectra were assigned to the magnetosome protein MamO, the top-10 proteins with highest SpC already covered 75% of all spectra (top-20 - 89%). Besides MamS and MamP, all identified genuine Mam and Mms magnetosome membrane proteins (according to Table 2) were also detected in the membrane shaving assay, however with highly variable SpC. Additionally, MamJ, MamA and MamD2 were also identified with comparably low SpC (suppl. Table S 4). Although with low SpC, only MGR_2730 and the MAI-encoded MGR_4114 were identified from the list of non-Mam/Mms proteins, but assigned genuine magnetosome proteins (Table 2). As in whole magnetosomes, MGR_3691 was again identified with a high SpC (suppl. Table S 4), emphasizing its role as a candidate magnetosome membrane protein. Six proteins previously not found in any of the analyzed fractions had assigned peptide spectra, of which MGR_1410, a predicted ammonia permease, had the highest SpC. Notably, also FeoB1 was detected with intermediate SpC. Most other identified proteins, most of them with a comparably low SpC, were components of transporters or of redox pathways and other proteins of the energymetabolism. The results from the Proteinase K membrane shaving assay again suggest that magnetosome membranes are specifically enriched mainly with Mam and Mms proteins.

606

607

608

609

610

611

612

613

614

615

616

617

618

619

620

621

622

623

624

625

626

628	Predictions of protein stoichiometry suggest that the magnetosome membrane is
629	densely packed with integral proteins.
630	Recently, based on quantitative Western blots with MamC-GFP labeled
631	magnetosomes, a conservatively estimated number of approximately 100 molecules
632	of MamC was suggested for an average sized magnetosome [21]. Based on this
633	number and the calculated $A_{mag.str(N)}/A_{mag.str(MamC)}$ -values, we estimated the putative
634	copy numbers of the integral genuine magnetosome proteins for an average wild type
635	magnetosome of 45.5 nm diameter [3] (Table 2). Within the membrane, TMHs of
636	integral proteins are associated with one or two boundary lipid shells that interact
637	with the hydrophobic protein domain. The diameter of the most prevalent lipid head
638	groups in magnetosome membrane is approximately 0.72 nm. Hence, an annular
639	boundary lipid shell would increase the diameter of an embedded TMH to 2.5 and
640	4.0 nm for one and two boundary lipid layers, respectively (Figure 3A and B). TMHs
641	of multi-membrane spanning proteins might be packed without internal lipid
642	boundary layers. According to Jacobson et al. [22], a diameter of 2.4 nm can be
643	assumed for the whole transmembrane domain (TMD) of a packed tetraspan-protein,
644	and a diameter of 3.2 nm for a packed heptaspan-protein (Figure 3C-E). We
645	interpolated these values for magnetosome membrane proteins that exhibit 1 to 18
646	TMHs and calculated the average TMH-coverage of the magnetosome membrane
647	(surface) based on the predicted copy numbers of the proteins and different boundary
648	lipid assumptions (see experimental procedures for details) (Table 4). We chose to
649	focus only on genuine Mam and Mms proteins, since the copy numbers of other
650	potential integral magnetosome-membrane proteins are negligible, together
651	accounting for only 2 % (Table 2).
652	According to this calculation, TMDs of magnetosome proteins already cover 18-20%
653	of the magnetosome surface, without taking into account boundary lipids. If one shell

of boundary lipid is added, this coverage increases to 62-97% and further to impossible 131-238% if a second shell of boundary lipids is included. Assuming a mixed TMH-packing model in reality, this indicates that TMHs of all proteins seem to be in close contact to each other and in most cases can be only surrounded by one layer of boundary lipids. We used the alternative PAI quantification method to estimate protein abundance and membrane coverage (suppl. Table S 5 and suppl. Table S 6, full calculation can be found in suppl. File 1). Using PAI, we even estimated higher copy numbers for most proteins and up to 170% higher membrane coverages (suppl. Table S 6). Therefore, the magnetosome membrane seems to be very rigid and tightly packed with trans-membrane proteins and only contains a smaller number of "free" lipids. For comparison, the hepta-spanning archaeal bacteriorhodopsin is one of the most tightly clustered transmembrane proteins and might be present in up to 600,000 molecules on a surface of 15 µm² [42]. Assuming a trimer configuration of the molecule with an approximate TMD-diameter of 5.2 nm [PDB] (including 3 enclosed lipids), this would indicate a total membrane coverage of 28%, which is somewhat higher but in the same range as our conservatively estimated coverage of the magnetosome membrane.

Conclusions

654

655

656

657

658

659

660

661

662

663

664

665

666

667

668

669

670

671

672

673

674

675

676

677

678

In summary, our proteomic data and modelling estimated the composition of the integral magnetosome-membrane proteome of MSR-1. Although the prediction of relative protein abundancies from label-free mass spectroscopy data is inherently error-prone and might lead to under- or overestimations for individual proteins, the cautious interpretation of our data allowed us to approximate the protein composition and coverage of the magnetosome membrane in a novel approach. Absolute quantifications of other individual magnetosomes-membrane proteins in the future

will further refine our model, which will be the basis for more precise determination of the structure of this unique bacterial organelle. By directly comparing protein abundances of magnetosomes purified from single magnetosome gene deletion mutants with those of the wild-type, our approach could be utilized to systematically assay the interdependency on protein localization to the organelle. This might prove as a powerful tool to further investigate the complex interaction-network of magnetosome proteins. Finally, our prediction of an unusually crowded protein composition within the membrane of the organelle also might substantially contribute to the assumption that a lipid raft like association of magnetosomemembrane proteins takes place prior to the magnetosome invagination [3,5]. Altogether our results will help to elucidate the processes involved in biogenesis of magnetosomes.

691 Acknowledgments

679

680

681

682

683

684

685

686

687

688

689

690

695

- This work was supported by grants from the Deutsche Forschungsgemeinschaft
- 693 (Schu1080/9-2 and 15-3) and the European Research Council (Proposal N° 692637
- 694 Syntomagx) to Dirk Schüler.

References

- 696 [1] F. Popp, J.P. Armitage, D. Schüler, Polarity of bacterial magnetotaxis is
- controlled by aerotaxis through a common sensory pathway, Nat. Commun. 5
- 698 (2014). doi:10.1038/ncomms6398.
- 699 [2] E. Cornejo, P. Subramanian, Z. Li, G.J. Jensen, Dynamic Remodeling of the
- Magnetosome Membrane Is Triggered by the Initiation of Biomineralization,
- 701 7 (2016) 1–9. doi:10.1128/mBio.01898-15.Editor.
- 702 [3] O. Raschdorf, Y. Forstner, I. Kolinko, R. Uebe, J.M. Plitzko, D. Schüler,

- Genetic and Ultrastructural Analysis Reveals the Key Players and Initial Steps
- of Bacterial Magnetosome Membrane Biogenesis, PLoS Genet. 12 (2016) 1-
- 705 23. doi:10.1371/journal.pgen.1006101.
- 706 [4] R. Uebe, D. Schüler, Magnetosome biogenesis in magnetotactic bacteria, Nat.
- 707 Rev. Microbiol. 14 (2016) 621–637. doi:10.1038/nrmicro.2016.99.
- 708 [5] H. Nudelman, R. Zarivach, Structure prediction of magnetosome-associated
- 709 proteins, Front. Microbiol. 5 (2014) 1–17. doi:10.3389/fmicb.2014.00009.
- 710 [6] Y.A. Gorby, T.J. Beveridge, R.P. Blakemore, Characterization of the bacterial
- 711 magnetosome membrane., J. Bacteriol. 170 (1988) 834-41.
- 712 http://jb.asm.org/content/170/2/834.
- 713 [7] K. Grünberg, C. Wawer, B.M. Tebo, D. Schüler, A Large Gene Cluster
- Fig. 714 Encoding Several Magnetosome Proteins Is Conserved in Different Species of
- 715 Magnetotactic Bacteria, Appl. Environ. Microbiol. 67 (2001) 4573–4582.
- 716 doi:10.1128/AEM.67.10.4573-4582.2001.
- 717 [8] M. Tanaka, Y. Okamura, A. Arakaki, T. Tanaka, H. Takeyama, T. Matsunaga,
- Origin of magnetosome membrane: proteomic analysis of magnetosome
- membrane and comparison with cytoplasmic membrane., Proteomics. 6 (2006)
- 720 5234–47. doi:10.1002/pmic.200500887.
- 721 [9] K. Grünberg, E.-C. Müller, A. Otto, R. Reszka, D. Linder, M. Kube, R.
- Reinhardt, D. Schüler, Biochemical and Proteomic Analysis of the
- Magnetosome Membrane in Magnetospirillum gryphiswaldense, Appl.
- 724 Environ. Microbiol. 70 (2004) 1040–1050. doi:10.1128/AEM.70.2.1040-
- 725 1050.2004.

- 726 [10] M. Richter, M. Kube, D.A. Bazylinski, T. Lombardot, F.O. Glöckner, R.
- Reinhardt, D. Schüler, Comparative genome analysis of four magnetotactic
- bacteria reveals a complex set of group-specific genes implicated in
- magnetosome biomineralization and function., J. Bacteriol. 189 (2007) 4899–
- 730 910. doi:10.1128/JB.00119-07.
- 731 [11] S. Ullrich, M. Kube, S. Schübbe, R. Reinhardt, D. Schüler, A Hypervariable
- 732 130-Kilobase Genomic Region of Magnetospirillum gryphiswaldense
- 733 Comprises a Magnetosome Island Which Undergoes Frequent
- Rearrangements during Stationary Growth, J. Bacteriol. 187 (2005) 7176–
- 735 7184. doi:10.1128/JB.187.21.7176.
- 736 [12] T. Matsunaga, M. Nemoto, A. Arakaki, M. Tanaka, Proteomic analysis of
- irregular, bullet-shaped magnetosomes in the sulphate-reducing magnetotactic
- bacterium Desulfovibrio magneticus RS-1., Proteomics. 9 (2009) 3341–3352.
- 739 doi:10.1002/pmic.200800881.
- 740 [13] A. Lohße, S. Ullrich, E. Katzmann, S. Borg, G. Wanner, M. Richter, B. Voigt,
- 741 T. Schweder, D. Schüler, Functional Analysis of the Magnetosome Island in
- Magnetospirillum gryphiswaldense: The mamAB Operon Is Sufficient for
- Magnetite Biomineralization, PLoS One. 6 (2011) e25561.
- 744 doi:10.1371/journal.pone.0025561.
- 745 [14] U. Heyen, D. Schüler, Growth and magnetosome formation by
- 746 microaerophilic Magnetospirillum strains in an oxygen-controlled fermentor.,
- 747 Appl. Microbiol. Biotechnol. 61 (2003) 536–44. doi:10.1007/s00253-002-
- 748 1219-x.
- 749 [15] D. Schüler, R. Uhl, E. Bäuerlein, A simple light scattering method to assay

- 750 magnetism in Magnetospirillum gryphiswaldense, FEMS Microbiol. Lett. 132
- 751 (1995) 139–145. doi:10.1016/0378-1097(95)00300-T.
- 752 [16] D. Schultheiss, D. Schüler, Development of a genetic system for
- Magnetospirillum gryphiswaldense., Arch. Microbiol. 179 (2003) 89–94.
- 754 doi:10.1007/s00203-002-0498-z.
- 755 [17] S. Ullrich, D. Schüler, Cre-lox-based method for generation of large deletions
- 756 within the genomic magnetosome island of Magnetospirillum
- 757 gryphiswaldense., Appl. Environ. Microbiol. 76 (2010) 2439–44.
- 758 doi:10.1128/AEM.02805-09.
- 759 [18] F. Bonn, J. Bartel, K. Büttner, M. Hecker, A. Otto, D. Becher, Picking
- vanished proteins from the void: How to collect and ship/share extremely
- dilute proteins in a reproducible and highly efficient manner, Anal. Chem. 86
- 762 (2014) 7421–7427. doi:10.1021/ac501189j.
- 763 [19] S. Wolff, H. Hahne, M. Hecker, D. Becher, Complementary analysis of the
- vegetative membrane proteome of the human pathogen Staphylococcus
- 765 aureus., Mol. Cell. Proteomics. 7 (2008) 1460–8. doi:10.1074/mcp.M700554-
- 766 MCP200.
- 767 [20] J.A. Vizcaíno, A. Csordas, N. del-Toro, J.A. Dianes, J. Griss, I. Lavidas, G.
- Mayer, Y. Perez-Riverol, F. Reisinger, T. Ternent, Q.-W. Xu, R. Wang, H.
- Hermjakob, 2016 update of the PRIDE database and its related tools, Nucleic
- 770 Acids Res. 44 (2016) D447–D456. doi:10.1093/nar/gkv1145.
- 771 [21] S. Borg, J. Hofmann, A. Pollithy, C. Lang, D. Schüler, New Vectors for
- 772 Chromosomal Integration Enable High-Level Constitutive or Inducible

- Magnetosome Expression of Fusion Proteins in Magnetospirillum
- gryphiswaldense., Appl. Environ. Microbiol. 80 (2014) 2609–16.
- 775 doi:10.1128/AEM.00192-14.
- 776 [22] K. Jacobson, O.G. Mouritsen, R.G.W. Anderson, Lipid rafts: at a crossroad
- between cell biology and physics., Nat. Cell Biol. 9 (2007) 7–14.
- 778 doi:10.1038/ncb0107-7.
- 779 [23] F.D. Gunstone, J.L. Harwood, F.B. Padley, The Lipid Handbook, second edi,
- 780 Chapman & Hall/CRC, London, 1994.
- 781 [24] J. Rappsilber, U. Ryder, A.I. Lamond, M. Mann, Large-Scale Proteomic
- Analysis of the Human Spliceosome, Genome Res. 12 (2002) 1231–1245.
- 783 doi:10.1101/gr.473902.
- 784 [25] O. Raschdorf, J.M. Plitzko, D. Schüler, F.D. Müller, A tailored galK
- 785 counterselection system for efficient markerless gene deletion and
- chromosomal tagging in Magnetospirillum gryphiswaldense, Appl. Environ.
- 787 Microbiol. 80 (2014) 4323–4330. doi:10.1128/AEM.00588-14.
- 788 [26] R. Arai, H. Ueda, A. Kitayama, N. Kamiya, T. Nagamune, Design of the
- linkers which effectively separate domains of a bifunctional fusion protein.,
- 790 Protein Eng. 14 (2001) 529–32.
- 791 http://www.ncbi.nlm.nih.gov/pubmed/11579220.
- 792 [27] R. Uebe, K. Junge, V. Henn, G. Poxleitner, E. Katzmann, J.M. Plitzko, R.
- 793 Zarivach, T. Kasama, G. Wanner, M. Pósfai, L. Böttger, B.F. Matzanke, D.
- 794 Schüler, The cation diffusion facilitator proteins MamB and MamM of
- Magnetospirillum gryphiswaldense have distinct and complex functions, and

- are involved in magnetite biomineralization and magnetosome membrane
- 797 assembly., Mol. Microbiol. 84 (2011) 818–835. doi:10.1111/j.1365-
- 798 2958.2011.07863.x.
- 799 [28] X. Wang, Q. Wang, W. Zhang, Y. Wang, L. Li, T. Wen, T. Zhang, Y. Zhang,
- J. Xu, J. Hu, S. Li, L. Liu, J. Liu, W. Jiang, J. Tian, Y. Li, L. Wang, J. Li,
- 801 Complete Genome Sequence of Magnetospirillum gryphiswaldense, Genome
- 802 Announc. 2 (2014) 2–3. doi:10.1128/genomeA.00171-14.Copyright.
- 803 [29] A. Scheffel, A. Gärdes, K. Grünberg, G. Wanner, D. Schüler, The major
- magnetosome proteins MamGFDC are not essential for magnetite
- biomineralization in Magnetospirillum gryphiswaldense but regulate the size
- of magnetosome crystals., J. Bacteriol. 190 (2008) 377-86.
- 807 doi:10.1128/JB.01371-07.
- 808 [30] D. Murat, A. Quinlan, H. Vali, A. Komeili, Comprehensive genetic dissection
- of the magnetosome gene island reveals the step-wise assembly of a
- prokaryotic organelle, Proc. Natl. Acad. Sci. U. S. A. 107 (2010) 5593–5598.
- 811 doi:10.1073/pnas.0914439107.
- 812 [31] A. Taoka, R. Asada, H. Sasaki, K. Anzawa, L.-F. Wu, Y. Fukumori, Spatial
- localizations of Mam22 and Mam12 in the magnetosomes of
- Magnetospirillum magnetotacticum., J. Bacteriol. 188 (2006) 3805–12.
- 815 doi:10.1128/JB.00020-06.
- 816 [32] N. Zeytuni, E. Ozyamak, K. Ben Harush, G. Davidov, M. Levin, Y. Gat, T.
- Moyal, A. Brik, A. Komeili, R. Zarivach, Self-recognition mechanism of
- MamA, a magnetosome-associated TPR-containing protein, promotes
- complex assembly., Proc. Natl. Acad. Sci. U. S. A. (2011).

- doi:10.1073/pnas.1103367108.
- 821 [33] D. Yamamoto, A. Taoka, T. Uchihashi, H. Sasaki, H. Watanabe, T. Ando, Y.
- Fukumori, Visualization and structural analysis of the bacterial magnetic
- organelle magnetosome using atomic force microscopy., Proc. Natl. Acad. Sci.
- 824 U. S. A. 107 (2010) 9382–7. doi:10.1073/pnas.1001870107.
- 825 [34] A. Lohße, S. Borg, O. Raschdorf, I. Kolinko, É. Tompa, M. Pósfai, D. Faivre,
- J. Baumgartner, D. Schülera, Genetic dissection of the mamAB and mms6
- operons reveals a gene set essential for magnetosome biogenesis in
- magnetospirillum gryphiswaldense, J. Bacteriol. 196 (2014) 2658–2669.
- 829 doi:10.1128/JB.01716-14.
- 830 [35] C. Rong, C. Zhang, Y. Zhang, L. Qi, J. Yang, G. Guan, Y. Li, J. Li, FeoB2
- Functions in Magnetosome Formation and Oxidative Stress Protection in
- Magnetospirillum gryphiswaldense Strain MSR-1., J. Bacteriol. 194 (2012)
- 833 3972–6. doi:10.1128/JB.00382-12.
- 834 [36] C. Lang, D. Schüler, Expression of green fluorescent protein fused to
- magnetosome proteins in microaerophilic magnetotactic bacteria., Appl.
- Environ. Microbiol. 74 (2008) 4944–53. doi:10.1128/AEM.00231-08.
- 837 [37] D. Murat, A. Quinlan, H. Vali, A. Komeili, Supporting Information -
- Comprehensive genetic dissection of the magnetosome gene island reveals the
- step-wise assembly of a prokaryotic organelle, Proc. Natl. Acad. Sci. U. S. A.
- 840 107 (2010).
- 841 [38] M. Tanaka, A. Arakaki, S.S. Staniland, T. Matsunaga, Simultaneously discrete
- biomineralization of magnetite and tellurium nanocrystals in magnetotactic

843 (2010)bacteria., Environ. Microbiol. 76 5526-32. Appl. 844 doi:10.1128/AEM.00589-10. 845 A. Poetsch, D. Wolters, Bacterial membrane proteomics, Proteomics. 8 (2008) 846 4100–4122. doi:10.1002/pmic.200800273. 847 Y. Okamura, H. Takeyama, T. Matsunaga, A Magnetosome-specific GTPase 848 from the Magnetic Bacterium Magnetospirillum magneticum AMB-1, J. Biol. 849 Chem. 276 (2001) 48183–48188. doi:10.1074/jbc.M106408200. 850 D. Schultheiss, R. Handrick, D. Jendrossek, M. Hanzlik, D. Schüler, The 851 Presumptive Magnetosome Protein Mms16 Is a Poly(3-Hydroxybutyrate) 852 Granule-Bound Protein (Phasin) in Magnetospirillum gryphiswaldense, J. Bacteriol. 187 (2005) 2416-2425. doi:10.1128/JB.187.7.2416. 853 854 [42] R.C.H. del Rosario, C. Oppawsky, J. Tittor, D. Oesterhelt, Modeling the 855 membrane potential generation of bacteriorhodopsin, Math. Biosci. 225 (2010) 856 68-80. doi:10.1016/j.mbs.2010.02.002.

858 Figures and tables

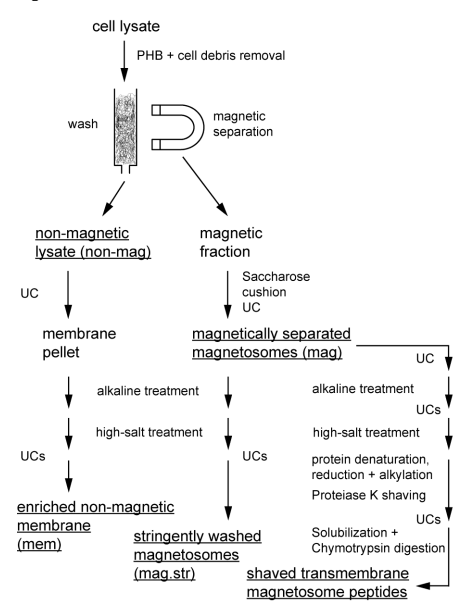


Figure 1: Fractionation workflow of magnetically separated magnetosomes, stringently washed magnetosomes, non-magnetic cell lysate and enriched non-magnetic membrane fraction. Additionally, workflow to obtain transmembrane peptides from magnetosome proteins by Proteinase K membrane shaving is outlined. UC: Ultracentrifugation

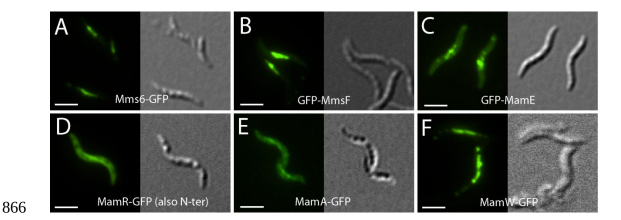


Figure 2: Cellular localization of six Mam/Mms proteins in *M. gryphiswaldense*. All proteins were C- or N-terminally fused to EGFP and expressed in the wild type. A 30 amino acid alpha-helical linker (HL) was placed in between the fusion proteins. A) C-terminal P_{mamDC}-mms6-HL-egfp construct, expressed from plasmid. B) N-terminal P_{mamDC}-mmsF-HL-egfp construct, expressed from plasmid. C) N-terminal P_{mamDC}-mamE-HL-egfp construct, expressed from ectopic chromosomal locus. D) C-terminal P_{mamDC}-mamR-HL-egfp construct, expressed from plasmid. N-terminal fusion exhibited comparable localization pattern. E) C-terminal mamA-HL-egfp construct, expressed from native chromosomal locus (in-frame gene fusion) F) C-terminal P_{mamDC}-mamW-HL-egfp construct, expressed from plasmid. Fluorescence (left) and corresponding differential interference contrast (right) images are shown. Scale bar: 2μm

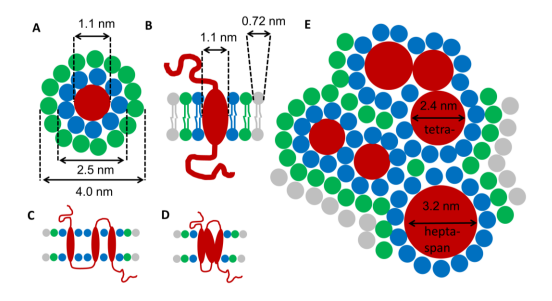


Figure 3: Model of transmembrane helices (domains) embedded in a membrane.

Transmembrane helices (domains) are colored in red, primary boundary lipids in blue, secondary boundary lipids in green and non-interacting ('free') lipids in grey. Approximate cross section diameters of several entities are indicated within the figure. A) Top view of single transmembrane helix with two layers of boundary lipids. B) Side view of transmembrane helix with two layers of boundary lipids and one layer of free lipids. C) Side view of three transmembrane helices of a single protein all embedded by boundary lipids. D) Side view of three transmembrane helices of a single protein packed in one domain embedded by boundary lipids but without internal lipids. E) Top view of an array of transmembrane domains of several proteins surrounded by variable numbers of boundary lipids. One of these proteins is a packed protein with transmembrane domains consisting of three transmembrane helices (tetra-span) and another protein with seven (hepta-span) helices.

Table 1: Number of assigned genuine magnetosome proteins under different

parameter conditions

Parameters									
$A_{\text{mem}}/A_{\text{non-mag}} \ge$	1	0	1	1	0	1	1	1	0
$A_{\rm mag}/A_{\rm mem}$ $>$	1	1	0	1	1	4	1	4	4
$A_{mag.str}/A_{mag} \ge$	1	1	1	0	0	1	1	1	1
$A_{\text{mag.str}}/A_{\text{mag.str}(MamC)}$ >	-	-	-	-	-	-	0.01	0.01	0.01
Total # of assigned proteins A	81	151	556	155	651	30	43	26	30
	81 23	151 24	556 24	155 24	651 27	30 21	43 23	26 21	30 22
Total # of assigned proteins A									

897

898 899

900

901

902

903

895

896

[encoded within mam or mms operons (including mamW, mamF2, mamD2 and ftsZm]

A total number of proteins that meet the applied parameter filter set (= assigned)

^B number of assigned proteins that are predicted magnetosome membrane proteins (MMPs)

^C number of proteins that are predicted MMPs, but do not meet parameter criteria

D number of assigned proteins that are not predicted MMPs

Table 2: Top20-abundant and all assigned genuine magnetosome proteins. The following parameters were applied: $A_{mem}/A_{non-mag} \geq 1$, $A_{mag}/A_{mem} > 4$, $A_{mag.str}/A_{mag} \geq 1$. Abundance relative to MamC was calculated by $A_{mag.str}/A_{mag.str(MamC)}$ value. The number of predicted protein copies is estimated by assuming a MamC copy number of 100 per magnetosome [21]. Genuine magnetosome-membrane proteins meeting the parameter criteria are marked by grey columns; other proteins only show high abundance values compared to MamC copy number.

Rank Abundance (A _{mag,str})	Abundance relative to MamC [predicted copy number]	Protein name [main putative function[4]]	Molecular weight
1	1.21 [121]	Mms6 [magnetite crystal nucleation and growth]	13 kDa
2	1.00 [100]	MamC [magnetite crystal growth]	12 kDa
3	0.51 [51]	MamD [magnetite crystal growth]	30 kDa
4	0.48	YajC [preprotein translocase subunit]	13 kDa
5	0.31 [31]	MamF2 [unknown]	12 kDa
6	0.31 [31]	MamE [magnetosome maturation, magnetite crystal nucleation]	78 kDa
7	0.26	MGR_0659 (Mms16) [Phasin]	16 kDa
8	0.25 [25]	MmsF [magnetite crystal growth]	14 kDa
9	0.24	MGR_3650 [Outer membrane protein (porin)]	41 kDa
10	0.23 [23]	MamB [magnetosomal iron transport, magnetosome membrane formation]	32 kDa
11	0.21 [21]	MamM [magnetosomal iron transport]	34 kDa
12	0.21 [21]	MamF [magnetite crystal growth]	12 kDa
13	0.20	MamR [magnetite crystal growth]	8 kDa
14	0.18	AtpF [ATP synthase B chain precursor]	19 kDa
15	0.18	MGR_1798 [Outer membrane protein]	17 kDa
16	0.17	MamA [magnetosome maturation]	24 kDa
17	0.17 [17]	MamY [magnetosome membrane maturation]	41 kDa
18	0.14 [14]	MamO [magnetite crystal nucleation]	65 kDa
19	0.13	AtpG [ATP synthase B' chain]	18 kDa
20	0.13	MamJ [magnetosome chain formation]	44 kDa
24	0.11 [11]	MamP [magnetite crystal nucleation and growth, redox control]	28 kDa
25	0.10 [10]	MamT [magnetosomal redox control]	19 kDa
28	0.09 [9]	MamS [magnetite crystal growth]	19 kDa
36	0.06 [6]	MamG [magnetite crystal growth]	8 kDa
40	0.06 [6]	MamI [magnetite crystal nucleation and growth]	8 kDa
43	0.05 [5]	MamW [unknown]	15 kDa
47	0.05 [5]	MGR_4114 [unknown]	7 kDa

58	0.04 [4]	MamH [unknown, putative iron importer]	46 kDa
60	0.04 [4]	MamN [magnetite crystal growth]	46 kDa
95	0.02 [2]	MGR_0622 [ATP-binding region, Histidine kinase]	50 kDa
100	0.02 [2]	MGR_2552 [Cytochrome oxidase maturation cbb3-type]	7 kDa
102	0.02 [2]	MGR_2833 [unknown]	26 kDa
111	0.02 [2]	MamX [magnetosomal redox control]	28 kDa
127	0.02 [2]	MamZ [magnetosomal redox control, putative iron importer]	72 kDa
187	0.01 [1]	MGR_2730 [unknown]	20 kDa
226	0.01 [1]	MGR_0916 [unknown]	9 kDa
444	0.00 [0]	MGR_0581 [unknown]	10 kDa
535	0.00 [0]	MGR_2491 [unknown]	8 kDa
1040	0.00 [0]	MGR_3321 [two-comp. sensor histidine kinase]	47 kDa

Table 3: Additional candidate genuine magnetosome membrane proteins predicted by less stringent filter parameters. Proteins here are only listed if not already mentioned in Table 2.

A _{mem} / A _{non-mag}	$ m A_{mag}/ m$	$ m A_{mag.str}/ m$	$A_{mag.str}/$ $A_{mag.str(MamC)}$	Rank (A _{mag.str})	$\begin{aligned} &\text{Set parameters: } A_{\text{mem}}/A_{\text{non-mag}} \geq 1, \\ &A_{\text{mag}}/A_{\text{mem}} > 1, A_{\text{mag,str}}/A_{\text{mag}} \geq 1, \\ &A_{\text{mag,str(N)}}/A_{\text{mag,str(MamC)}} \geq 0.02 \end{aligned}$
4.3	2.0	1.3	0.07	32	MamQ [magnetosome membrane maturation] (30 kDa)
3.4	1.3	1.5	0.07	34	MGR_3120 Bacterial sec-independent translocation protein mttA/Hcf106 (8 kDa)
2.9	1.3	1.3	0.07	35	MGR_1712 translocase, subunit Tim44 (26 kDa)
1.2	1.4	1.0	0.05	46	MGR_0255 conserved hypothetical protein (11 kDa)
5.0	3.7	1.3	0.03	66	MGR_1199 Peptidase M48, Ste24p (33 kDa)
1.1	1.2	1.2	0.02	87	MGR_4238 regulatory protein (22 kDa)
3.9	1.1	1.1	0.02	93	MGR_0007 glycosyl transferase, group 2 family (27 kDa)
6.6	1.3	1.9	0.02	114	MGR_0417 serine O-acetyltransferase (27 kDa)
4.3	1.9	1.1	0.02	129	MGR_3354 phos.lipid/glycerol acyltransferase (30 kDa)
A _{mem} / A _{non-mag}	A_{mag}/A_{mem}	A _{mag.str} / A _{mag}	$\begin{array}{l} A_{mag.str}/\\ A_{mag.str(MamC)} \end{array}$	Rank (A _{mag.str})	Set parameters: $A_{\text{mem}}/A_{\text{non-mag}} \ge 0$ $A_{\text{mag}}/A_{\text{mem}} > 4$, $A_{\text{mag.str}}/A_{\text{mag}} \ge 1$, $A_{\text{mag.str(N)}}/A_{\text{mag.str(MamC)}} > 0.01$
0.0	∞	1.1	0.20	13	MamR [magnetite crystal growth] (8 kDa)
0.4	6.5	1.5	0.10	27	MGR_2633 Phasin (12 kDa)
0.1	6.8	1.2	0.02	116	MGR_2416 cytochrome c (12 kDa)
0.8	5.0	1.1	0.01	142	MGR_1351 CreA (17 kDa)

Table 4: Coverage of the magnetosome membrane surface by transmembrane domains of integral magnetosome proteins

Integral magnetosome membrane coverage by TMDs ^A	All TMHs ^B are isolated from each other	TMHs of individual proteins are tightly packed
No boundary lipid	18 %	20 %
One boundary lipid	97 %	63 %
Two boundary lipids	238 %	132 %

920 A: Transmembrane domain

921 B: Transmembrane helix

918

922 Supplementary information

- 923 suppl. File 1: Calculations and interactive table to analyze magnetosome membrane
- 924 proteom data
- 925 Fig S 1: Organization of genes associated with magnetosome formation within the
- genomic magnetosome island (MAI) of M. gryphiswaldense.
- 927 Fig S 2: 2D SDS-PAGE of fractions employed for mass spectrometry analysis:
- 928 Fig S 3: Representative Transmission Electron Micrographs of M. gryphiswaldense
- wild type (A), $\Delta mgr3691$ ($\Delta MM22$) (B) and $\Delta mgr4114op$ (C). Scale bar represents
- 930 500 nm
- 931 suppl. Table S 1. Number of assigned genuine magnetosome proteins under different
- 932 parameter conditions
- 933 suppl. Table S 2: Number of genuine magnetosome proteins assigned by increasing
- 934 A_{mag}/A_{mem} ratios
- 935 suppl. Table S 3: Introduction of MamC abundance threshold to regulate the number
- 936 of assigned genuine magnetosome membrane proteins
- 937 suppl. Table S 4: Proteins with identified peptides from shaving assay.
- 938 suppl. Table S 5: Estimated number of protein copies for membrane integral
- 939 Mam/Mms Proteins according to different quantification methods.
- suppl. Table S 6: Coverage of the magnetosome membrane surface by
- transmembrane domains of integral magnetosome proteins calculated according to
- 942 different quantification methods
- 943 suppl. Table S 7: Strains and plasmids used in this study
- suppl. Table S 8: Oligonucleotides used in this study