A virus capsid-like nanocompartment that stores iron and protects bacteria from oxidative stress

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Abstract

Living cells compartmentalize materials and enzymatic reactions to increase metabolic efficiency. While eukaryotes use membrane-bound organelles, bacteria and archaea rely primarily on protein-bound nanocompartments. Encapsulins constitute a class of nanocompartments widespread in bacteria and archaea whose functions have hitherto been unclear. Here, we characterize the encapsulin nanocompartment from Myxococcus xanthus, which consists of a shell protein (EncA, 32.5 kDa) and three internal proteins (EncB, 17 kDa; EncC, 13 kDa; EncD, 11 kDa). Using cryo-electron microscopy, we determined that EncA self-assembles into an icosahedral shell 32 nm in diameter (26 nm internal diameter), built from 180 subunits with the fold first observed in bacteriophage HK97 capsid. The internal proteins, of which EncB and EncC have ferritin-like domains, attach to its inner surface. Native nanocompartments have dense iron-rich cores. Functionally, they resemble ferritins, cage-like domains, attach to its inner surface. Native nanocompartments have dense iron-rich cores. Functionally, they resemble ferritins, cage-like iron storage proteins, but with a massively greater capacity (~30,000 iron atoms versus ~3,000 in ferritin). Physiological data reveal that few nanocompartments are assembled during vegetative growth, but they increase fivefold upon starvation, protecting cells from oxidative stress through iron sequestration.

Keywords cryo-electron microscopy; encapsulin; ferritin; HK97 fold; oxidative stress

Subject Categories Metabolism; Microbiology; Virology & Host Pathogen Interaction

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Introduction

Compartmentation allows cells to confine enzymes to increase metabolic efficiency, to protect them from proteolysis or other challenges, and to sequester toxic products (Diekmann & Pereira-Leal, 2013). In bacteria and archaea, cytosolic compartments are mainly protein-bound (Kerfeld et al., 2010; Yeates et al., 2008, 2011). One such class is encapsulin nanocompartments whose common feature is that their shell protein encapsulin (formerly referred to as linocin-like protein) has an HK97-like fold (Sutter et al., 2008). This fold was first observed in the capsid protein of HK97, a lambdoid bacteriophage with a triangulation number (T-number) of 7 (Wikoff et al., 2000) (see below, Discussion). Since its discovery, the HK97 fold has been observed in the capsid proteins of several other tailed bacteriophages (Agirrezabala et al., 2007; Fokine et al., 2005; Jiang et al., 2006; Parent et al., 2010; Spilman et al., 2011; White et al., 2012), and cryo-EM data strongly suggest that it is also present in herpesvirus capsids (Baker et al., 2005; Cardone et al., 2012; Homa et al., 2013). Recently, it was found by X-ray crystallography that an archaeal encapsulin and a bacterial encapsulin have similar folds arrayed, respectively, in $T = 3$ (180-subunit) and $T = 1$ (60-subunit) icosahedral shells (Akita et al., 2007; Sutter et al., 2008). On this basis, it has been proposed that encapsulins and the capsid proteins of tailed phages share a common evolutionary origin (Abrescia et al., 2012; Akita et al., 2007; Forrer & Prangishvili, 2009; Heinemann et al., 2011; Sutter et al., 2008), a hypothesis supported by the presence of genes encoding phage-like proteins (e.g., a replicative helicase) nearby an archaeal encapsulin gene (Heinemann et al., 2011). However, viral capsids and encapsulin nanocompartments have different roles: While the former transport viral genomes from one cell to another, the latter are involved in metabolism. Importantly, despite the identification of encapsulin proteins in a variety of bacteria, including medically important species such as Mycobacterium tuberculosis (Rosenkranz et al., 1998), the in vivo functions of encapsulin nanocompartments have not yet been demonstrated.

Myxococcus xanthus is a social bacterium characterized by a complex developmental life cycle. Actively growing cells are motile and, upon amino acid starvation, aggregate to form elaborate spore-filled fruiting bodies (Zusman et al., 2007). These
developmental processes make *M. xanthus* an ideal model organism to study cellular adaptation to environmental signals. While studying vegetatively growing *M. xanthus* cells, we noted the production of round, 32-nm-diameter, seemingly proteinaceous particles. Although these particles were sparse, amino acid starvation resulted in a fivefold increase, enabling purification of sufficient quantities for biochemical and structural characterization. In this study, we have characterized these particles: In particular, they contain a dense core composed mainly of iron and phosphorus. Using a variety of electron microscopic techniques, we compiled a detailed account of the molecular architecture of their encapsulin shell and of the composition and organization of its dense core. Exploring the functional correlates of these particles, we found that in disruption and deletion mutants of the shell-forming protein EncA, whereby the iron-sequestering ability is lost, more cells die when exposed to oxidative stress in the form of H$_2$O$_2$ compared to wild-type cells. Taken together, these observations lead to a model for the iron-sequestering encapsulin nanocompartments of *M. xanthus* and support the hypothesis that their function involves the regulated uptake of iron to protect the cells from oxidative stress and its subsequent release when needed by iron-requiring cytosolic enzymes.

### Results

#### Protein composition of native *M. xanthus* encapsulin nanocompartments

Electron microscopy of negatively stained specimens confirmed that purified material comprised a homogeneous population of spherical particles, ~32 nm in diameter (Fig 1A). SDS–PAGE (Fig 1B) showed that the particles are composed of four proteins, which were identified by mass spectrometry and named EncA (MXAN_3556), EncB (MXAN_3557), EncC (MXAN_4464), and EncD (MXAN_2410). Gel quantitation yielded the estimate that EncA accounts for ~72% of the total protein content and the minor proteins EncB, EncC, and EncD for ~7,~15, and ~6%, respectively. Taken together, the minor proteins amount to ~40% of the EncA mass. Since each nanocompartment has 180 copies of EncA (see below), this is accompanied by, on average, ~36 copies of EncB, ~92 of EncC, and ~47 of EncD. Edman degradation analysis indicated that three of the proteins were incorrectly annotated in the NCBI database: EncA is seven aa shorter, EncB is six aa shorter, and EncC is fourteen aa longer.

Sequence analysis showed that EncA shares 24% sequence identity and 43% sequence similarity with the encapsulin shell protein

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**Figure 1.** Native encapsulin nanocompartments are composed of four proteins and contain a dense iron-rich core.

A. EM of negatively stained nanocompartments. Scale bar, 100 nm.
B. SDS–PAGE of purified native encapsulin nanocompartments. Mass markers are indicated (in kDa).
C. Domain maps of the four encapsulin proteins. Yellow, HK97-like domain; orange, ruberythrin/ferritin-like domain; asterisks, ExxH metal coordination motifs; gray, encapsulation signal.
D. EM of unstained/air-dried nanocompartments. Scale bar, 100 nm.
E. STEM and energy-dispersive X-ray spectroscopy (EDX) images of two representative particles. In the STEM image (unstained specimen), positive signals from mass scattering identify two nanocompartments, while the corresponding EDX images map elemental concentrations of Fe and P (strongly above background), and S and Ca (at or close to background). Red crosshairs indicate identical positions on one nanocompartment. Scale bar, 100 nm.

Source data are available online for this figure.
of *Thermotoga maritima* (Fig 1C and Supplementary Fig S1). Furthermore, EncB, EncC, and EncD were found each to have a C-terminal motif (LTVGSLLR; Fig 1C and Supplementary Figs S2, S3 and S4) similar to those proposed to serve as encapsulation signals in other systems (Fan *et al*., 2010; Sutter *et al*., 2008). No other protein encoded in the *M. xanthus* genome (Goldman *et al*., 2006) contains this motif. Using the Pfam server (Finn *et al*., 2014), we found that EncB and EncC—but not EncD—possess rubrerythrin/ferritin-like domains. Bioinformatic analysis of EncB and EncC shows that in both proteins, this domain is highly conserved, including their copies of the ExxH motif (Fig 1C and Supplementary Figs S2 and S3), which is known to bind iron and other metals (Liu & Theil, 2004; Sieker *et al*., 2000).

**Native nanocompartments have dense cores rich in iron and phosphorus**

During purification, we observed that encapsulin particles sediment at an unusually high density in cesium chloride gradients. Moreover, they have an electron-dense core that is conspicuous in electron micrographs of unstained preparations (Fig 1D). This core is significantly smaller—~24 nm versus ~32 nm—than the protein shell. To investigate its elemental composition, we analyzed these particles by energy-dispersive X-ray spectroscopy and found that their core is highly enriched for iron and phosphorus (Fig 1E and Supplementary Fig S5). No other element was detected in comparable amounts. To estimate the relative abundances of iron and phosphorus, similar preparations were also analyzed by inductively coupled plasma mass spectrometry (ICP-MS) (Table 1) which revealed a molar ratio of approximately 4:1 for Fe:Pi.

**Native encapsulin nanocompartments and recombinant EncA shells are *T* = 3 icosahedra**

Visualized by cryo-EM, native nanocompartments from *M. xanthus* appear as round 32-nm-diameter particles (Fig 2A). These images confirmed that a large majority—~94% in this preparation—have electron-dense cores (a minority particle lacking such a core is marked with a black arrow in Fig 2A). When EncA was expressed recombinantly in *E. coli*, most (~80%) of the particles produced were of the same size as native nanocompartments, that is, ~32 nm (Fig 2B), while ~20% were smaller, mostly ~18 nm in diameter (e.g., white arrow in Fig 2B). These observations implied that EncA is capable of self-assembly, but with somewhat lower fidelity in size determination than native *M. xanthus* particles.

To obtain a high-resolution reconstruction of the shell, we focused on the 32-nm recombinant EncA shells. A data set of ~14,000 images yielded a density map at 4.6 Å resolution that

<table>
<thead>
<tr>
<th>Sample</th>
<th>Iron (µg/l)</th>
<th>Phosphorus (µg/l)</th>
<th>Fe:Pi ratio</th>
</tr>
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<tbody>
<tr>
<td>Control</td>
<td>73.40 ± 75.20 (102)</td>
<td>&lt; 1 LOD</td>
<td></td>
</tr>
<tr>
<td>Encapsulin</td>
<td>276.600 ± 7.764 (2.81)</td>
<td>72.390 ± 1.495 (2.07)</td>
<td>4.1</td>
</tr>
<tr>
<td>Ferritin</td>
<td>268.200 ± 21.370 (7.94)</td>
<td>30.160 ± 72.6 (2.41)</td>
<td>9.1</td>
</tr>
</tbody>
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Values for each element are the average concentration plus or minus the standard deviation in µg/l, calculated based on at least 5 experimental replicates per sample. All protein samples were dissolved in deionized water, which was also used as control solution. Note that the measured Fe:Pi ratio of the used horse spleen ferritin is in agreement with published data (Michaelis *et al*., 1943). Parentheses, coefficient of variation, in % (standard deviation divided by the average concentration for each sample and multiplied by 100).

**Figure 2. Cryo-EM of encapsulin nanocompartments and EncA shells.**

A, B Cryo-micrographs of (A) native nanocompartments and (B) recombinantly expressed EncA capsids.

C Section through a reconstruction of the EncA capsid. An enlargement of the boxed area showing a longitudinal section through an α-helix is inset (upper right).

D Atomic model of the T = 3 EncA capsid. The three quasi-equivalent subunits are colored yellow, green, and red. Symmetry axes are marked.

E The 3-subunit asymmetric unit fitted into electron density (transparent).

F, G Atomic model of EncA, rainbow-colored with the N-terminus in blue, and the C-terminus in red.

Data information: Scale bars: (A, B) 50 nm; (C) 10 nm; (D) 5 nm; (E, C insert) 2 nm.
depicts a shell with icosahedral symmetry and a triangulation number of $T = 3$ (180-subunits) (Fig 2C–E and Supplementary Fig S6). In it, $\alpha$-helices are patent in slices through the map (Fig 2C), and landmark features of the HK97 fold (Wikoff et al., 2000), including the “spine” $\alpha$-helix, the A- and P-domains, and the $\beta$-hairpin E-loop, are evident. Accordingly, its information content was considered sufficient to justify the construction of an atomic model (Fig 2D–G, Supplementary Figs S7 and S8 and Supplementary Table S2). To do so, we first generated a homology model using the encapsulin protein from *T. maritima* (pdb 3DKT) as template and docked it as a rigid body into the EM map. This structure was then adapted by “flexible fitting” (Trabuco et al., 2008), whereby the starting model was systematically modified to optimize the fit. Finally, the structure was refined using X-ray crystallographic procedures, applying constraints that compensate in part for the limited resolution (Adams et al., 2010).

Starting with other references, for example, the encapsulin protein from *Pyrococcus furiosus* (pdb 2E0Z), led to essentially the same end-product.

The dense cores of native nanocompartments consist of multiple 5-nm granules

From the similar diameters and scalloped peripheries of both the 32-nm EncA shells from *E. coli* and the dense-cored encapsulin particles from *M. xanthus*, we expected the latter to have an icosahedral shell similar to that of the EncA shell. However, the native particles posed a challenge for reconstruction in that the images are dominated by the core material, which is not icosahedrally ordered, as we initially inferred from visual inspection and subsequently confirmed by cryo-ET (see below). This property made it difficult to determine the particles’ orientations, a necessary step for calculating a three-dimensional reconstruction. This problem was solved by limiting the correlated areas used for orienting the particles to a peripheral annulus to which the core material does not contribute (Fig 3A). The fact that only a small fraction (~10%) of the image area was used restricted the information content, leaving greater uncertainties in the Euler angles used to define orientations; in turn, this limited the resolution. Nevertheless, this approach yielded a
reconstruction at a resolution of 17 Å (Fig 3B and C): It depicts an icosahedral $T=3$ surface shell whose outer surface is essentially indistinguishable from that of the EncA capsid when the latter is limited to the same resolution. The reconstruction shows the internal contents as three layers of density of which the densest is the middle one, a 3-nm-thick shell peaking at a radius of 6 nm. However, this reconstruction conveys the core structure after it has been subjected to extensive inter-particle averaging and icosahedral symmetrization. In order to obtain 3-dimensional structures of individual particles that had not been symmetrized, we imaged them by cryo-ET (Fig 4A–C), which yielded density maps with a resolution of 40–50 Å. The tomograms revealed that the core is not a continuous structure but consists of discrete granules. The number of granules per particle ranges from 11 to 19, averaging 14. They are typically round and 5–6 nm in diameter (Fig 4D). The granules are not regularly distributed—in particular, neither in number nor in their spatial distribution do they comply with icosahedral symmetry. Most of the granules are offset from the capsid, with the commonest offset (~5 nm) matching the radial position of the densest layer in the icosahedral reconstruction (Fig 4B and C). The combined volume of granules within a single nanocompartment is variable but averages ~1,600 nm³.

The minor proteins line the interior surface of the EncA shell

A minor fraction (~6%) of the particles isolated from *M. xanthus* lack dense cores (black arrow in Fig 2A). To investigate whether they nevertheless contain the putative internal proteins (EncB, EncC and EncD), we calculated a cryo-EM reconstruction (Fig 5A). Although the resolution was limited to 24 Å by their paucity ($N = 370$), it does indeed show internal densities underlying the EncA shell (Fig 5A). These internal densities are most evident at the three-fold axes, and they are not present in a similarly calculated reconstruction of the EncA shell (Fig 5B). The locations and shapes of these densities are better seen in a difference map (Fig 5C and D; and Supplementary Fig S9). However, they are less dense than the shell, indicative of partial occupancy. We interpret these data to mean that these proteins bind sub-stoichiometrically to defined sites on the inner surface of the capsid.

The dense cores contain on average ~30,000 iron atoms

To measure the amount of cargo in the native encapsulin particles, we imaged them by STEM, a technique in which the signal recorded at each pixel is proportional to the mass locally sampled by the

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**Figure 4. Cryo-ET of encapsulin nanocompartments with electron-dense cores.**

A Tomographic slice showing several native *Myxococcus xanthus* encapsulin nanocompartments.

B Tomographic slices through two nanocompartments (top and bottom rows).

C Gallery of tomographic central sections of eight nanocompartments.

D Histogram of the diameters of electron-dense granules in the nanocompartment cores.

Data information: Scale bars, 25 nm.
Iron storage in encapsulin nanocompartments

Figure 5. Cryo-EM visualization of internal proteins.

A–C Central sections through reconstructions of (A) native nanocompartments lacking a dense core; (B) EncA capsids; (C) the difference map between (A) and (B). In (C), contours delineating the outer and inner surfaces of the EncA capsid are superimposed in half of the section to aid in locating densities of internal proteins. White arrows label densities in direct contact with the inner surface, and black arrows label offset densities.

D Isosurface rendering of a cutaway view of the EncA capsid (green-blue) and the positive difference density (orange) attributable to the internal proteins. Scale bar, 10 nm.

electron probe (Goldsbury et al., 2011; Thomas et al., 1994) (Fig 6). These experiments were carried out with a preparation that was essentially devoid of empty particles, as ascertained by cryo-EM (data not shown). All of the particles had electron-dense cores, albeit of variable sizes (Fig 6A and B). Their masses give a symmetric unimodal distribution with a mean of 10.0 MDa and a standard deviation (SD; σ) of 0.74 MDa (Fig 6C). The SD reflects both genuine particle-to-particle variability (σ₁) and the spread (σ₂) to be expected in a set of STEM mass measurements from homogeneous protein particles of the same size. As these sources of variability are independent, they are related by σ² = σ₁² + σ₂². We estimated σ₂ from the SD of a set of mass measurements made on 56-nm segments of tobacco mosaic virus (TMV), which contain approximately the same amount of protein as the encapsulin nanocompartments. The TMV virions have a known and uniform mass density (131.4 kDa/nm) and serve as mass calibrators. Present on the same grid as the encapsulin particles, they were prepared and imaged under identical conditions. These data gave 7.4 ± 0.43 MDa (Fig 6D); accordingly, σ₁ = 0.60 MDa.

We used these data to calculate the mean and range of the cargo mass and its payload of iron atoms. Since an EncA monomer has a mass of 32.5 kDa, that of the shell (180 subunits) is 5.85 MDa. We estimated the amount of internal proteins (EncB, EncC, and EncD) (see above and Fig 1B) to be 40% of EncA, that is, 2.34 MDa. This estimate is consistent with one obtained from the 3D reconstruction of iron-free native nanocompartments (Fig 5A) in which the number of voxels in the shell is compared with that occupied by the internal proteins. It follows that the mass of dense material (total particle mass minus combined mass of the EncA shell and internal proteins) is, on average, 1.81 MDa. The SD calculated above gives a range (mean ± σ₁) of 1.21 to 2.41 MDa.

If we infer from the ICP-MS data (see above) that iron and phosphorus are the only significant elemental components and they are present in a molar ratio of 4 (Fe; 56 Da) to 1 (P; 31 Da), then the average nanocompartment contains ~28,500 iron atoms. If the phosphorus is in the form of phosphate (PO₄, 95 Da), this number falls to ~22,500. However, for particles at the upper end of the mass range (mass = mean + σ₁), these numbers increase to ~38,000 and ~30,000 iron atoms, respectively. Given the residual ambiguity in this calculation, we take 30,000 as a round number for the average payload of iron atoms. In any case, this payload is an order of magnitude larger that that of ferritins, which has been estimated to average ~3,000 iron atoms (Andrews, 2010).

Encapsulin nanocompartment assembly is induced under starvation conditions

A genetic study of M. xanthus found the EncA gene to be required for fruiting body formation, though it was not recognized as an encapsulin homolog at the time (Kim et al., 2009). Fruiting body formation is a developmental program triggered by starvation and resulting in sporulation (Zusman et al., 2007). Since starvation leads to increased assembly of loaded nanocompartments (see above), we measured the production of EncA protein and of encapsulin nanocompartments during starvation and sporulation. Actively growing and starved cells contained comparable amounts of EncA (Fig 7A). However, isolation of nanocompartments from starved cells yielded a fivefold increase in yield, compared to actively growing cells (Fig 7B). The enhanced production of nanocompartments by starved cells was confirmed by direct visualization of the iron cores of assembled particles in EM of unstained thin sections (Fig 7C). This suggests that encapsulin production is regulated not only at the level of gene expression, but also post-translationally at the level of assembly. Neither nanocompartments nor EncA were found at significant levels in mature spores after 48 or 96 h of development (Fig 7A and B, and Supplementary Fig S10), raising the possibility that pre-existing nanocompartments were eliminated by proteolytic degradation to recycle their iron and phosphorus contents.

Encapsulin nanocompartments protect cells from oxidative stress caused by peroxide

Fruiting body formation is accompanied by a massive die-off of cells whereby ~80% of vegetative cells lyse and a small proportion proceed to sporulation (Lee et al., 2012a; Wireman & Dworkin, 1977). Since iron can catalyze the production of toxic reactive oxygen species through the Fenton reaction (Cornelis et al., 2011), the assembly of encapsulin nanocompartments during starvation may serve a protective role against cellular death due to oxidative
stress by reducing the amount of iron free in the cytosol. To test this hypothesis, we investigated the effects of mutations in EncA (Supplementary Fig S11) on the survival of *M. xanthus* under oxidative stress caused by peroxide (H$_2$O$_2$) exposure. Both the independently generated disruption mutant *encA::pCR2.1* and the in-frame deletion mutant Δ*encA* showed increased sensitivity to 0.5 mM H$_2$O$_2$ confirming that these structures are indeed important for oxidative stress survival (Fig 7D and E).

**Discussion**

**Encapsulin nanocompartments encapsulate iron**

Iron is essential to life because it serves as a cofactor of many enzymes. However, it can also produce highly reactive hydroxyl radicals that cause cellular damage (Harrison & Arosio, 1996). To regulate the supply of free intracellular iron, cells have evolved protein assemblies that act as iron storage containers. The best known are the three subfamilies of ferritins: classical ferritins, bacterioferritins, and “DNA-binding proteins from starved cells” (Dps proteins), of which classical ferritins and bacterioferritins function in iron storage and Dps proteins in iron detoxification (Andrews, 2010; Theil *et al*, 2013). Ferritins are multimeric complexes whose subunits, each a four-helix bundle, assemble into hollow shells with cubic symmetry (Zeth, 2012). These “molecular cages” can accommodate substantial quantities of iron (Andrews, 2010). The shells are penetrated by channels through which iron ions can enter to be stored inside the complex and subsequently released under iron-requiring conditions.

Encapsulin nanocompartments appear to be operationally similar to ferritins, despite marked differences in size (32 nm versus 11 nm), storage capacity for iron—which is greater by an order of magnitude, shell symmetry (icosahedral vs. octahedral), the fold of the building block (HK97-like versus a 4-helix bundle), and in that the iron deposits nucleate on cargo proteins rather than directly on to the shell protein.

Although there is, to our knowledge, no prior history of studies into the occurrence of ferritins in *M. xanthus*, genome analysis reveals that it encodes several putative ferritin-like proteins (MXAN_2878, MXAN_2342, MXAN_1008, and Dps proteins MXAN_0866 and MXAN_1562) besides those packaged into encapsulin particles. Pending further information, it appears plausible that *M. xanthus* may use its ferritins for iron homeostasis during vegetative growth and, upon starvation, induce a second iron storage system that functions independently alongside the ferritin system. The second (encapsulin-based) system would allow the cells to temporarily sequester iron and phosphorus as an antioxidant response or to be used later for nutritional purposes (Theil, 2011).

**Comparison of *M. xanthus* encapsulin with other encapsulins and with HK97 gp5***

A growing body of evidence indicates that encapsulin proteins generically possess HK97-like folds, as do the capsid proteins of
other tailed bacteriophages as well as herpesviruses, often in the complete absence of sequence similarity. This mutual resemblance is apparent on comparing the encapsulin structures of *M. xanthus*, *T. maritima*, and *P. furiosus* (Fig 8A–D). Differences between EncA and the other encapsulins are observed primarily in the E-loop (expected, as these are flexible regions) and in the insertion loop anterior to the spine helix (which is absent in *P. furiosus*). Of note, these three molecules have an N-terminal
α-helix (blue in Fig 8A–C) that is absent from HK97 but present in the capsid proteins of some other bacteriophages (e.g., T4 (Fokine et al., 2005), P22 (Parent et al., 2010), and 80x (Spilman et al., 2011)). Since this helix faces the interior of the encapsulin shell, it might be involved in interactions with the internal proteins. On comparing EncA with HK97 gp5* (Fig 8A, D and E), the most pronounced differences, other than the absence of the N-terminal α-helix in gp5*, are in the A-domains where EncA has an extra α-helix (yellow in Fig 8A), and in the E-loops and the tips of the P-domains which differ slightly.

Two distinctive features of HK97 gp5 are: (i) that it undergoes maturational proteolysis in which the N-terminal Δ-domain is excised as gp5 (41 kDa) is converted to gp5* (31 kDa) (Duda et al., 1995); and (ii) the maturing capsid undergoes a sequence of large-scale conformational changes en route from the first assembled precursor Prohead I to the fully mature Head II (reviewed by (Hendrix & Johnson, 2012; Steven et al., 2005)). To what extent do encapsulins share these properties? Here, we restrict the discussion to M. xanthus encapsulin.

The Δ-domain appears to serve as a scaffolding protein that guides assembly (Huang et al., 2011a). Essentially, all phage capsids
with T-numbers of 7 or greater employ a Δ-domain in this role, either appended as an N-terminal fusion (as in HK97) or as a separate gene product. The morphologically correct particles generated upon expressing EncA in E. coli indicate that this protein does not need a scaffolding protein. The reason may reflect the lower T-number (3), there being numerous examples of T = 3 capsids that self-assemble efficiently without a scaffolding protein, although few if any examples have been documented for viral capsids with the HK97 fold. In this context, the 20% of smaller particles among the recombinant expression products in E. coli (see above) are ~18 nm in diameter and conform to T = 1 (60-subunit) icosahedral symmetry (unpublished results).

The conformational state of EncA shells resembles that of HK97 in its immature form

The successive states of the HK97 maturation pathway are distinguished primarily by rigid-body rotations of the subunits as the intermolecular interactions are shifted to different surfaces (Conway et al., 2001), although there are also some local adjustments (Gertsman et al., 2009). Which state of the maturing HK97 capsid does the encapsulin shell most closely resemble? We addressed this question in two ways: by comparing the fold of EncA monomer with the various HK97 gp5* conformers and by comparing the respective capsomers. These calculations placed the EncA monomer marginally closer to the Head II gp5* monomer (the RMSDs obtained were 2.17 Å for Head II and 2.76 Å for Prohead II; alignment was performed using the superpose program from CCP4 (Collaborative Computational Project, 1994)) (compare Fig 8A with D and E). On the other hand, EncA capsomers are clearly most similar to those of Prohead II (Fig 8F), although the EncA hexamers are not skewed (those of Prohead II are skewed, resembling two dislocated trimers (Conway et al., 1995)). Of note, we did not find any evidence of a similar expansion process in EncA, and such a transition has yet to be reported for any other encapsulin. We speculate that this may reflect the absence of an evolutionary mandate to increase the encapsulin shell’s capacity for cargo and to confer additional stability, as applies to HK97 and other phage capsids.

A functional model of the encapsulin nanocompartment

Taken together, current observations led us to the nanocompartment model shown in Fig 9. It envisages substoichiometric binding of EncB, EncC, and EncD to specific sites (of which there are 180) on the inner surface of an EncA shell via their common C-terminal binding motif. We anticipate that iron atoms may enter the shell via narrow channels through the capsid wall, of which those located on the fivefold axes or between the A-domain and the spine helix of adjacent subunits from the same capsomer (Fig 9B) are the most likely candidates. Entering iron atoms are then incorporated into nascent granules nucleated on and around the metal-binding motifs on the rubrerythrin domains of EncB and EncC. Known structures for rubrerythrin domains depict a bundle of four lengthy α-helices. As the corresponding sequences in EncB and EncC are only long enough to accommodate two such helices (Supplementary Figs S2 and S3), it is plausible that they dimerize. Between the rubrerythrin domains and the C-terminal binding motifs, there are tracts that may afford flexible linkers long enough to account for the observed, non-icosahedrally symmetric distribution of iron-containing granules. As the internal proteins are reasonably well ordered and shell associated in iron-free shells (Fig 5), it appears likely that the buildup of iron deposits on the EncB and EncC proteins is accompanied by detachment from the EncA shell.
Granulation of the iron-containing core of native *M. xanthus* nanocompartments as opposed to the formation of a single mineralized nugget may be advantageous in terms of rapidity of uptake and subsequent release. If mineralization initiates at 15 sites (the average number of granules) instead of one, iron uptake can proceed proportionately more rapidly. Conversely, if there are 15 granules of 5.5 nm in diameter (taken to be spherical for the sake of this discussion) instead of the same amount of iron packed in a single supergranule of 13.5 nm in diameter, the amount of surface area exposed, from which iron atoms may presumably be shed, is almost three times greater (~1,425 nm² versus 572 nm²).

**A diversity of encapsulin functions, natural and synthetic**

Only recently has it been recognized that the cytoplasms of many bacteria are populated with protein-bound nanocompartments (Kerfeld *et al.*, 2010; Yeates *et al.*, 2008, 2011). One such class is the encapsulins whose shell proteins possess a fold previously found only in viral capsids. However, unlike virus capsids, which serve as vehicles for genome transport, encapsulins are involved in metabolism. To date, only a few have been identified on the basis of structural similarity but we anticipate that this set will grow considerably. One subset of encapsulin operons also code for a dye-decolorizing peroxidase (DyP) or for a ferritin-like protein (Flp), which was taken to suggest a role in the oxidative-stress response (Sutter *et al.*, 2008), now amply confirmed in the *M. xanthus* system. However, other encapsulin nanocompartments have been suggested to perform different functions, including antibacterial activity in *Brevibacterium linens* (Valdes-Stauber & Scherer, 1994), proteolytic activity in *T. maritima* (Hicks *et al.*, 1998), and lignin degradation activity in *Rhodococcus jostii* RHA1 (Rahmanpour & Bugg, 2013).

The encapsulation signal at the C-termini of EncB, EncC, and EncD suggests the possibility of appending this short amino acid sequence to other exogenous proteins for the purpose of engineering novel encapsulin particles housing a variety of activities with a broad range of applications ranging from imaging to therapeutics (Huang *et al.*, 2011b; Lee *et al.*, 2012b; Theil *et al.*, 2013). Moreover, close scrutiny of the differences between HK97 gp5, which assembles into 48 nm $T = 7$ shells, may make it possible to adapt EncA in order to assemble EncA nanocompartments larger than the native 32 nm $T = 3$ form.

### Materials and Methods

#### Strains and media

The wild-type strain was *M. xanthus* DK1622 (Kaiser, 1979). Vegetative cells were grown at 32°C in rich CTT medium (1% casitone, 1 mM K$_2$PO$_4$, 8 mM MgSO$_4$, 10 mM Tris/HCl, final pH 7.6) or in starvation TPM medium (1 mM K$_2$PO$_4$, 8 mM MgSO$_4$, 10 mM Tris/HCl, final pH 7.6) (Hodgkin & Kaiser, 1979). For preparation of agar plates and maintenance of cells on solid medium, 1.5% agar was added to CTT before autoclaving.

The *encaA*::pCR2.1 (EH410) disruption strain was constructed by recombination of the suicide plasmid pCR2.1 containing a 396-bp homology region into the MXAN_3556 (*enca*) gene. The in-frame deletion strain Δenca (EH411) was constructed using an allelic exchange by homologous recombination strategy, with plasmid pBJ113 as described previously (Ueki *et al.*, 1996). Both procedures are described in Supplementary Materials and Methods.

#### Large-scale purification

DK1622 cells grown in CTT medium to an Abs$_{550}$ of 0.2 were transferred into TPM medium and grown for 18–24 h to induce nanocompartment assembly. Cells were harvested, re-suspended in water or Tris buffer, chilled, and broken by ultrasonication. Cellular debris was removed by pelleting at 16,000 g and again after adding 15% (w/v) ammonium sulfate. Polyethylene glycol was added to the supernatant and incubated on ice for 45 min. Precipitated proteins were harvested at 47,800 g, re-suspended in water or Tris buffer, and CsCl added to a concentration of 0.3 g/ml. Ten milliliters of this solution was layered over a 2-ml cushion of 5.7 M CsCl and spun for 16 h at 35,000 rpm in a Beckman SW55 rotor. The nanocompartments formed a whitish band near the top of the CsCl cushion and were harvested and stored at 4°C. For more concentrated preparations, the sample was dialyzed against water or buffer to remove the CsCl, then layered over a 10–40% sucrose gradient, and centrifuged for 2.5 h at 20,000 rpm. Nanocompartment-containing fractions were combined and re-centrifuged for 1.5 h at 55,000 rpm in a Beckman W55 rotor. The nanocompartments formed a small brownish pellet that could easily be re-dissolved in water or buffer and was stored at 4°C. Negative staining EM and SDS–PAGE were used to confirm the purity of the preparations.

#### Identification of protein components of native encapsulin nanocompartments

The proteins in several independent isolates were identified by two approaches: MALDI/TOF mass spectrometry and Edman degradation analysis (Supplementary Table S1). Purified nanocompartments at ~0.1 mg/ml protein were run on a SDS–PAGE gel from which bands were excised and analyzed by MALDI/TOF mass spectrometry by Protea Biosciences, Inc. (Morgantown, West Virginia) or the Johns Hopkins University High Throughput Center Mass Spectrometry Facility. To identify the N-termini, bands separated on SDS–PAGE were blotted onto PVDF membranes, stained with colloidal Coomassie and subjected to Edman degradation at the Biochemistry Core Facility of the Max Planck Institute of Biochemistry ( Martinsried, Germany).

#### Inductively coupled plasma mass spectrometry

Purified nanocompartments were dissolved in reverse osmosis-purified water, which was also used as control. Analysis was performed on samples extracted with 1% HNO$_3$ and 0.5% HCl, using an Agilent 7500ce ICP-MS. Elemental standards were used to determine the limit of detection for each element (0.066 g/l for iron and 6.77 μg/l for phosphorus). Final measurements consisted of 5 independent isolations done in triplicate for the nanocompartments, water control, and 150 μl of a horse spleen ferritin solution of 0.8 mg/ml as internal standard (Fitzgerald Industries International).
Energy dispersive X-ray analysis

Samples consisting of either purified nanocompartments or adenovirus (strain Ad5.CMV-GFP 10MR2000, Quantum) were applied to carbon-coated copper grids. STEM and EDX data were acquired at the High-Resolution Analytical Electron Microbeam Facility of the Integrated Imaging Center in the Departments of Earth and Planetary Sciences and Biology of Johns Hopkins University using a Philips CM 300 FEG microscope operating at 300 kV equipped with an Oxford light element energy dispersive X-ray spectroscopy detector. Gatan DigitalMicrograph and ES Vision software packages were used for subsequent analyses. Iron was consistently found at high levels in the core of the encapsulin particles and was not present in the proteinaceous rim of the particles (the EncA shell) nor in background areas.

Cloning of the encA gene in E. coli and preparation of EncA capsids

The expression vector was constructed by amplifying the coding region of the encA (MXAN_3556) gene and cloning this fragment into plasmid pET151-d/TOPO (Invitrogen), thus generating plasmid pods. For expression, E. coli BL21 cells harboring the plasmid were grown to log phase in LB medium and then vigorously shaken for 45 min, after which precipitated proteins were pelleted. The pellets were resuspended in Tris-Triton X buffer (10 mM Tris pH 7.6 and 10 mM MgCl₂, 0.1% Triton X-100), and cesium chloride added to give a density of 1.3 g/ml. Tubes were centrifuged for 16 h at 35,000 rpm. Capsids formed a whitish band near the top of the gradient, which was recovered and analyzed by SDS-PAGE and Western blot and checked by negative stain EM. To remove detergent for cryo-EM, samples were diluted in 2 ml of 10 mM Tris pH 7.6 and 10 mM MgCl₂ and centrifuged for 3 h at 360,000 × g. The pellet was re-suspended in 30 µl of Tris buffer and stored at 4°C.

Cryo-electron microscopy and image reconstruction

Purified nanocompartments or EncA shells were applied to glow-discharged Quantifoil grids bearing a thin carbon film, blotted, vitrified, and imaged with a CM200 FEG microscope, as described (Cheng et al., 1999). For high-resolution analysis, the same preparation was imaged on an FEI Polara at 59,000× magnification. Micrographs recorded on Kodak SO163 film at 1- to 2-µm (CM200) or 0.5- to 1.6-µm (Polara) defocus were digitized at 1.23 Å/pixel (CM200) or 1.10 Å/pixel (Polara). Image processing was performed with the Bsoft package (Heymann & Belnap, 2007), as described (Nemecek et al., 2012). Images were CTF corrected by phase flipping in the initial stages and by phase and baseline compensation in later stages. Particles were picked manually, and origins and orientations were determined by projection matching. To obtain an unbiased initial reference, four sets, each of ~200 EncA capsids, were reconstructed, initially assigning random orientations (Yan et al., 2007). The resulting density maps were quite consistent and were further validated by the variance-based procedure VIVA (Cantele et al., 2003). One of these maps was used as initial reference, leading to a reconstruction at ~8 Å resolution from ~8,000 CM200 images. The hand of the reconstruction was established by the tilt-pair method (Belnap et al., 1997; Cheng et al., 2002). For higher resolution, ~14,000 particles from Polara micrographs were reconstructed in similar fashion, using the CM200 EncA map filtered to 15 Å as initial reference. It had a resolution of 4.6 Å according to the 0.143 FSC criterion (Scheres & Chen, 2012). Fourier amplitudes were “sharpened” by using the P. furiosus crystal structure as reference.

To reconstruct dense-cored nanocompartments, only the rims of the shell were used for particle alignment and orientation determination. The initial reference was generated as described above. The final reconstruction, combining information from ~900 particles, had ~17 Å resolution, at 0.3 FSC cutoff.

To visualize the internal proteins, a reconstruction of the EncA shell was subtracted from a reconstruction of native nanocompartments lacking electron-dense cores. The latter reconstruction, calculated from ~400 images, had a resolution of ~25 Å at 0.3 FSC cutoff. The former reconstruction was calculated using ~400 randomly selected EncA images for a resolution of ~23 Å. A difference map calculated after scaling the two maps consistently, using a purpose-built algorithm (Heymann et al., 2003). To estimate the cumulative mass of internal proteins, the reconstruction of native nanocompartments lacking electron-dense cores was segmented into two regions, the shell and the internal content, using Bsoft, and the number of voxels corresponding to each region were integrated using MAPMAN.

Atomic modeling of EncA

An initial homology model was generated using I-TASSER (Zhang, 2008) with the T. maritima encapsulin as reference (pdb 3DKT). This model was flexibly fitted into the EM map, using MDDF (Trabuco et al., 2008). The resulting atomic coordinates were then used to generate the three quasi-equivalent subunits of an asymmetric unit by rigid-body fitting into the corresponding densities and propagated over the icosahedral shell, using Chimera (Pettersen et al., 2004). Additionally, the EM density map was placed in a cubic cell using Bsoft, MAPMAN (Kleywegt & Jones, 1996), and MAPMASK tools (Collaborative Computational Project, 1994). Structure factors and Hendrickson–Lattman coefficients, describing phases, were calculated using the phenix.map_to_structure_factors program from PHENIX (Adams et al., 2010), and the atomic model was refined with phased maximum-likelihood target as implemented in phenoX.refine (Afonine et al., 2005). To compensate for the limited resolution, icosahedral restraints, Ramachandran restraints, and weighting of the stereochemical restraints were employed. Manual model building in Coot (Emsley et al., 2010) alternated with automatic refinement. Refinement statistics are shown in Supplementary Table S2. R-free is not reported. If calculated, the R-free would be similar to the R-value because of the high non-crystallographic symmetry (Kleywegt & Brunger, 1996).

Cryo-electron tomography

Grids bearing thin films of native nanocompartment suspension were mixed with 10-nm colloidal gold particles coated with bovine serum albumin (BSA) (Aurion, Wageningen, the Netherlands) to serve as fiducial markers and vitrified as described above.
Single-axis tilt series were recorded on a Tecnai-12 TEM (FEI) equipped with a GIF 2002 energy filter (Gatan, Warrendale, PA) operated in zero-loss mode with an energy slit width of 20 eV. Images were acquired on a 2048 × 2048 pixel CCD camera using SerialEM (Mastronarde, 2005) at a magnification of 38,500× (0.78 nm/pixel) and 4-μm nominal defocus. The tilt range was from −66° to +66° with images recorded every 2°, for a total electron dose of ~70 e-/Å². According to the NLOO-2D method (Cardone et al., 2005), the in-plane resolution of the tomograms was 40–50 Å.

Scanning transmission electron microscopy

Samples of nanocompartments mixed with TMV were adsorbed for 1 min to a titanium grid bearing a thick holey carbon film, washed 10 times with water, and then blotted and plunged into liquid nitrogen slush. Freeze-drying, grid transfer into the microscope, and imaging were performed as previously described (Sen et al., 2007). Dark-field micrographs were recorded at raster steps of 1.0 or 2.0 nm per pixel. Masses for individual nanocompartments and TMV segments were calculated using PCMass (available from the Brookhaven STEM resource). The nanocompartment preparation used for this analysis was different than the one used for cryo-EM/cryo-ET. Both preparations were found to be consistent by negative staining EM and SDS–PAGE except for the fact that the cryo-EM/cryo-ET preparation contained ~6% of the core-lacking particles while the STEM one contained < 1%.

Quantitative analysis of encapsulin expression and assembly

For Western blotting, 1 × 10⁸ cells grown in CTT and TPM medium and 1 × 10⁸ purified spores were collected by centrifugation. Cells were lysed by adding 30 μl of SDS–PAGE loading buffer to this pellet and boiling for 10 min. Spores were re-suspended in 30 μl loading buffer and broken by ultrasonication, before boiling. Breakage was confirmed by light microscopy, using a Zeiss Axiosmager. The lysates were loaded onto a 4–20% Tris-Glycine gel (Invitrogen) and transferred to a membrane. The primary antibody, G3576, was generated using a standard immunization protocol for rabbit against the KLH-conjugated target peptide RLRGESGVVVSTGR, and the secondary antibody (goat anti-rabbit) was from Jackson ImmunoResearch Laboratories Inc. Reactive bands were visualized with Pierce ECL Western Blotting Chemiluminescent Substrate (Thermo Scientific).

To quantify yields of dense-cored nanocompartments, equal samples of the respective isolates were applied for 15 s to carbon-coated 400 mesh copper grids and the numbers of particles per area counted. At least three independent fields were analyzed in each experiment. Means and standard deviations were determined, and all values normalized against the count of nanocompartments from CTT-grown cells. The total number of particles counted was 735 for wild-type cells grown in CTT, and 3,899 for wild-type cells grown in TPM.

Isolation of spores

Spots containing ~4 × 10⁹ wild-type cells/ml were placed on TPM agar and incubated for 48 or 96 h to induce fruiting body formation and spore development. Fruiting bodies were harvested using a cell scraper and re-suspended in Tris buffer (10 mM Tris/HCl pH 7.5). Vegetative cells were disrupted by ultrasonication and heating (60 min 50°C), and spores were concentrated by centrifugation (10 min, 14,000 × g). To remove more cellular debris, the spores were re-centrifuged using a 15% (w/v) Percoll gradient (GE Healthcare Sciences). Upon centrifugation (30 min 47,800 × g), membranes and cellular debris floated on top, while the spores formed a pellet at the bottom of the tube. After removal of the supernatant, the spores were washed twice with buffer, before being centrifuged and stored at 4°C.

Embedding of cells for thin sectioning

Vegetative cells and spores were briefly centrifuged and then high-pressure-frozen, using a Leica EM HPM100 system. Frozen cells were transferred into pure acetone containing 2% glutaraldehyde and cryo-substituted at ~−80°C. After 72 h, the samples were gradually warmed and embedded in low-viscosity Spurr medium. Polymerized resin blocks were trimmed and thin sectioned using a diamond knife (Diatome). Silver-to-gold sections were mounted on copper grids and viewed without staining on a CM120 Philips electron microscope. Images were taken with a 4 × 4 ORCA ER CCD camera (Hamamatsu).

Oxidative stress survival assays

Cells were grown in rich medium and then treated with hydrogen peroxide. A protocol similar to that used to assay hydrogen peroxide stress survival in M. tuberculosis was performed, with modifications described (Wu et al., 1997; Lu et al., 2010). DK1622 and encA cells were grown to mid-log phase in CTT medium and then diluted to an Abs₅₅₀ of 0.2 units with fresh CTT. Next, 10-ml samples of each culture were transferred to sterile 25-ml glass flasks (Fisher Scientific) and were either left untreated or treated with a final concentration of 0.5 mM hydrogen peroxide, with each treatment done in triplicate. Cultures were incubated at 32°C with shaking at 250 rpm. Cell survival was monitored by measuring the Abs₅₅₀ of each sample at 0, 1, 3, and 6 h after the addition of peroxide, and the values were normalized and expressed as a percentage of the starting absorbance in each tube. As described in Supplementary Materials and Methods, the disruption and deletion strains used in these experiments were generated completely independently.

Data deposition

The EncA refined coordinates were deposited in the protein data bank (PDB ID 4PT2). The cryo-EM maps of the T = 3 EncA shells and of the core-lacking M. xanthus encapsulin particles were deposited in the Electron Microscopy Data Bank (accession codes EMD-5917 and EMD-5953).

Supplementary information for this article is available online: http://emboj.embopress.org

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Author contributions
CAM, JF, DN, ACS, and EH designed research; CAM, JF, DN, NC, AAA, DW, ASL, JSW, and EH performed research; [BH contributed to analytic tools; CAM, JF, DN, ACS, and EH analyzed data; and JF, CAM, EH, and ACS wrote the paper with contributions from all other authors.

Conflict of interest
The authors declare that they have no conflict of interest.

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