The Vault Exterior Shell Is a Dynamic Structure that Allows Incorporation of Vault-Associated Proteins into Its Interior†

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Received May 26, 2006; Revised Manuscript Received August 3, 2006

ABSTRACT: Vaults are 13 million Da ribonucleoprotein particles with a highly conserved structure. Expression and assembly by multimerization of an estimated 96 copies of a single protein, termed the major vault protein (MVP), is sufficient to form the minimal structure and entire exterior shell of the barrel-shaped vault particle. Multiple copies of two additional proteins, VPARP and TEP1, and a small untranslated vault RNA are also associated with vaults. We used the SF9 insect cell expression system to form MVP-only recombinant vaults and performed a series of protein-mixing experiments to test whether this particle shell is able to exclude exogenous proteins from interacting with the vault interior. Surprisingly, we found that VPARP and TEP1 are able to incorporate into vaults even after the formation of the MVP vault particle shell is complete. Electrospay molecular mobility analysis and spectroscopic studies of vault-interacting proteins were used to confirm this result. Our results demonstrate that the protein shell of the recombinant vault particle is a dynamic structure and suggest a possible mechanism for in vivo assembly of vault-interacting proteins into preformed vaults. Finally, this study suggests that the vault interior may functionally interact with the cellular milieu.

Vaults are structurally conserved ribonucleoprotein (RNP)1 particles that have been implicated in multidrug resistance, in nucleocytoplasmic transport, and as scaffolds for both epidermal growth factor signaling and interferon-γ-activated JAK/STAT signaling pathways; however, their precise function remains unclear (1–9). The particles have a capped-barrel morphology with dimensions of approximately 41 nm × 41 nm × 72.5 nm (10) and, at 13 million Da, are the largest known RNP. Vaults have 8-fold symmetry around their longitudinal axis, and each half-vault appears to be identical (8-2-2 symmetry). When plated onto poly-L-lysine-coated electron microscopy grids and visualized by freeze-etch platinum shadowing, vaults tend to open and flatten into two half-vaults resembling flowerlike structures with eight petals symmetrically arranged around a central ring (11). Vaults are formed from three proteins: the 96 kDa major vault protein (MVP) which accounts for more than 70% of the particle mass, 290 kDa telomerase-associated protein 1 (TEP1), and the 193 kDa vault poly(ADP-ribose) polymerase (VPARP). The RNA component of vaults is an untranslated small RNA, termed vault RNA (VR) (1, 12–14). Expression of MVP alone in insect cells using a MVP-encoding baculovirus leads to the assembly of recombinant MVP-only vaults (henceforth termed MVP recombinant vaults), demonstrating that multimerization of this single protein is sufficient to form the exterior shell of vaults (15). In cells, VPARP and TEP1 are also found in nonvault cytoplasmic and nuclear fractions, and although both proteins have been shown to associate with telomerase activity, neither is required (14, 16–18). TEP1 is a RNA binding protein that is required for the stable association of VR with vaults and binds to VR and telomerase RNA in an indirect yeast three-hybrid assay and directly in electrophoretic mobility shift assays (17, 19, 20). VPARP is a functional PARP family member that ADP-ribosylates itself and MVP and is the first vault component demonstrated to have an enzymatic activity (13). The precise stoichiometry of VPARP, TEP1, and VR with respect to a single vault particle is not yet known, nor is it clear whether all vaults contain equal numbers of these vault-associated components, as other cellular factors may influence their ability to assemble into vaults. Furthermore,
some species such as humans and bullfrogs express multiple related VRs that can interact with vaults (6, 12, 21).

The three-dimensional image reconstruction of rat vaults at 22 Å resolution using cryo-electron microscopy (cryoEM) revealed that they consist of a protein shell, 2–5 nm thick, surrounding a large interior space (22, 23). Individual vaults contain interior mass as seen in cryoEM images or TEM of negatively stained samples; however, these contents are either variable for a given vault or irregular in structure or position and thus are subtracted out during reconstruction (23). Image reconstruction combined with difference mapping is a powerful tool for determining subunit locations. Difference mapping of untreated and RNase-treated vaults purified from rat liver localized VR to the interior of the vault caps (22). Recombinant vaults formed in insect cells have the same structure as native vaults. Cysteine peptide-tagged MVP (cpMVP) recombinant vaults were resolved to 16 Å, the best resolution vault structure obtained to date (10). Difference mapping of vaults comprised of N-terminal epitope-tagged MVP with those made from untagged MVP revealed that the N-terminus of each MVP monomer lies at the waist of the vault particle (10). Co-infection of insect cells with baculoviruses encoding MVP and either TEP1, VPARP, or both leads to the formation of recombinant vaults containing these proteins (10). In addition, when a MVP-interaction domain (INT, the C-terminus of VPARP, amino acids 1471–1724) is fused to heterologous proteins, either luciferase or a green fluorescent protein variant, green lantern (GL), the fusion proteins have been shown to assemble into MVP recombinant vaults (13, 24). CryoEM reconstructions indicate that all of the vault-interacting components are bound to the inner surface of the vault particle (10, 24).

The large interior volume of the vault (spacious enough to encompass an intact ribosome) supports the hypothesis that the function of vaults is related to scaffolding, sequestration, or transport. However, nothing is known about how MVP monomers might assemble to form the exterior shell of vaults or how the other vault components assemble into the vault interior. Although none of the vault components are covalently associated, virtually all of the MVP in cells biochemically fractionates with the vault complex (1, 6). Following purification, vault particles are highly stable, able to resist dissociation after treatment with 2 M urea, 1.0% Triton X-100, or pH extremes ranging from 4 to 10 (11). Thus, it seems unlikely that the MVP shell of vaults disassembles into subunits and then reassembles to allow entry of other proteins and/or contents. It is possible that vault contents are packaged simultaneously with the assembly of the vault MVP shell. Another non-mutually exclusive hypothesis is that intact vaults could open and close in response to cellular signals or that the vault shell in vivo is elastic, allowing relatively free exchange with the cytoplasmic environment. This latter possibility is reminiscent of findings that some viral capsids in solution are dynamic structures in which polypeptides and nucleic acids, mapped by X-ray crystallography as internal, are significantly more susceptible to external agents than would be expected (25–27).

In this study, we show that the vault particle is a dynamic structure allowing the vault-associated proteins VPARP and TEP1 to form complexes with recombinant vaults (expressed using the Sf9 insect cell system) even after the MVP shell of vaults has formed. These experiments were carried out by mixing vault-interacting proteins with preformed MVP recombinant vaults. In addition, we show that the vault-interacting proteins localize to the interior of the intact vault particle. Our results suggest that recombinant vault particles exhibit substantial flexibility that imparts access to the interior volume. The dynamic nature displayed by the recombinant vaults used in this study likely represents a key mechanism for the interaction of endogenous vaults with their in vivo environment.

### EXPERIMENTAL PROCEDURES

**Micrococcal Nuclease Digestion of Purified Vaults and Subcellular Extracts.** Rat liver vaults (4 μg) were purified as previously described (1). Purified vaults or phenol-extracted RNA from purified vaults was treated with 2000 units/mL micrococcal nuclease for 0, 15, 30, or 60 min at 37 °C. Crude vault extracts (100000 g pellet, P100) were prepared from normal rat kidney epithelial cells (LR117) as previously described (6). The P100 pellet was resuspended by dounce homogenization in buffer [50 mM Tris-HCl (pH 7.4), 1.5 mM MgCl2, and 75 mM NaCl] containing 10% glycerol, 1 mM phenylmethylsulfonyl fluoride, and protease inhibitors. DEproteinized RNA was isolated by phenol/chloroform extraction and ethanol precipitation. Equal volume amounts of P100 (RNP) or deproteinized P100 (RNA) extracts were treated with 2000 units/mL micrococcal nuclease for 20 min at 25 °C. All micrococcal nuclease digestions were terminated by the addition of EGTA to a final concentration of 8 mM. Reaction mixtures were extracted with phenol and chloroform, precipitated with ethanol, fractionated on a 10% acrylamide gel, and electrophoretically transferred to Zeta GT+ membrane (Bio-Rad), and probed for VR as previously described (6).

**Plasmids and M-INT Cloning and Expression.** Construction of the VPARP, TEP1, GL-INT, and CP-MVP (cysteine-rich N-terminal epitope tag-MVP)-expressing baculoviruses is described elsewhere (10, 24). The C-terminal MVP-interaction domain of VPARP (amino acids 1563–1724, M-INT) was inserted into Escherichia coli expression vector pET28a (Novagen). M-INT was expressed in *E. coli* strain BL21 CodonPlus (Stratagene) and purified on a His-bind column (Novagen). The eluted protein was dialed into buffer [20 mM Tris (pH 7.9), 50 mM KCl, 0.1 mM EDTA, and 10% glycerol] containing 1 mM DTT and 1 mM PMSF. Purified protein was aliquoted and stored at −80 °C.

**Sf9 Insect Cells and Vault Purifications.** Cells were maintained in SF-900 II SFM media and grown at 27 °C. Cultures were infected with MVP, TEP1, VPARP, or GL-INT viruses at a multiplicity of infection (MOI) of 0.01 for approximately 65 h and then pelleted and lysed on ice in buffer A [50 mM Tris-HCl (pH 7.4), 75 mM NaCl, and 0.5 mM MgCl2] with 1% Triton X-100, 1 mM dithiothreitol, 0.5 mM PMSF, and protease inhibitor cocktail (Sigma). Vault purification from lysates was performed as previously described (15). In experiments involving the mixing of lysates from two separate infections, each cell pellet was lysed separately and then the two were mixed and incubated together on ice for 30 min before proceeding with the standard vault purification. Pellets from the 20–60% sucrose fractions were each resuspended in 20 mM MES buffer (pH 6.5) for further analysis.
Gas-Phase Electrophoretic Mobility Molecular Analyzer (GEMMA). The details of the GEMMA have been described elsewhere (28–30). Briefly, the GEMMA (TSI Inc., St. Paul, MN) consists of an electrospray ionization (ESI) unit with a neutralizing chamber, a differential mobility analyzer (DMA), and a condensation particle counter (CPC). The DMA voltage was scanned, and data were recorded by Aerosol Instrument Manager Software. The software corrects for the efficiency of the condensation particle counter as well as for a small fraction of multiply charged particles assuming the bipolar charge distribution of Fuchs (31) and calculated by Wiedensohler’s formula (32). The electrophoretic mobility diameter maxima for peaks of interest were calculated from this corrected data. Igor Pro 4.08 software was used for further processing, including smoothing of spectra.

From the GEMMA measurements by our laboratory (unpublished data) and from those reported by Bacher et al. (28) for more than 40 protein complexes ranging in size from small protein dimers to complexes as large as the 700 kDa 70S proteasome and megadalton-range viral particles, an average density of approximately 0.6 g/cm³ can be used to calculate the molecular mass of small and large protein complexes. Data presented for the molecular mass of vault particles are means ± the standard error.

Prior to analysis, samples were desalted using Millipore Microcon centrifugal filter devices (YM-100) in 20 mM ammonium acetate (pH 7.4). Concentrations of analyses ranged from 400 to 900 ng/μL. Protein solutions were introduced via the ESI source at a flow rate of approximately 70 nL/min. The electrophoretic mobilities for the samples used to estimate molecular masses were determined from three independent experiments with MVP vaults and MVP–VPARP vaults from lysate mixing and from two independent experiments with MVP–VPARP vaults from co-infection, in which the total protein concentration desalted and analyzed ranged from 40 to 600 ng/μL.

Negative Staining and Electron Microscopy. Vault-containing samples were prepared for electron microscopy by absorption of samples onto a fresh 1.8 nm thick carbon film mounted on carbon-coated holey-film grids for 5 min at 4 °C. Following sample adsorption, grids were floated for 5 min on 1 mL of 1% uranyl acetate at 4 °C and dried on filter paper prior to being viewed in a JEM 1200EX microscope. The carbon-coated holey-film grids and carbon film were prepared by S. Ryazantsev (Department of Biological Chemistry, David Geffen School of Medicine). Micrographs were captured with a BioScan 600W digital camera (Gatan Inc., Pleasanton, CA) using Gatan’s DigitalMicrograph version 3.7.1. Images were further processed using Adobe Photoshop 6.0.

Green Fluorescence Quenching. Quenching of purified GL-INT vaults and measurement of fluorescence intensity were carried out as previously described (24).

In Vitro Binding and Thrombin Cleavage. Purified vaults were incubated with 20, 40, or 100 ng of M-INT for 1 h at 25 °C in buffer A with 1% Triton X-100 and 1 mM DTT in a 20 μL volume containing 1 mg/mL BSA. Volumes were then adjusted to 70 μL in the same buffer, and samples were pelleted at 60 000 rpm in a Beckman TLA 100.1 benchtop ultracentrifuge. Pellets were resuspended in 1× SDS sample buffer, and all samples were loaded onto 12% SDS—PAGE gels and transferred to Hybond-C (Amersham Biosciences) for Western blotting using polyclonal antibodies to MVP and VPARP. For cleavage reactions, 20 ng of M-INT was incubated with or without 1 μg of purified CP-MVP vaults for 1 h at 4 °C as described above. The volume was then adjusted to 55 μL, and a 10 μL aliquot was taken for the “zero” time point. Five microliters of thrombin (0.0004 unit/μL) or thrombin–agarose beads (0.01 unit/μL) was added, and the reaction mixture was incubated at 25 °C. Aliquots (10 μL) were taken out at 1 and 2 h and analyzed by Western blotting. Thrombin–agarose beads were prepared using biotinylated thrombin and streptavidin–agarose beads as described in the manufacturer’s instructions (Novagen). The beads were resuspended in buffer prior to use for the experiment described above.

RESULTS

Vault Particles Differ in Susceptibility to Nuclease Treatment Depending Upon Their Level of Purity. It has been previously demonstrated that the VR component of purified rat liver vaults can be removed by digestion with a cocktail of RNase A and RNase T1, despite difference maps of three-dimensional cryoEM reconstructions of RNase-treated and untreated vaults that localize VR to the presumably protected interior of the vault caps (22). Complete VR removal typically requires high concentrations of RNases and extended incubations for efficient digestion, supporting the notion that either the MVP vault shell is partially protective or the vault-associated proteins (VPARP and/or TEP1) impede the ability of RNases to access the interior of the vault caps. To study the accessibility of VR in the vault particle complex, we used micrococcal nuclease digestion. Micrococcal nuclease is a calcium-dependent nuclease that nonspecifically cleaves both RNA and DNA, and unlike RNases, time course measurements of digestion can be carried out, by chelating the calcium with EGTA to stop the reaction. When purified rat liver vaults were treated with 2000 units/mL micrococcal nuclease for up to 1 h at 37 °C, VR was substantially protected from digestion (Figure 1A, RNP). However, deproteinized VR extracted from rat liver vaults was susceptible to digestion under similar conditions (Figure 1A, RNA). Vault-containing fractions were then isolated from rat kidney epithelial cells by centrifugation of cell lysates at 100000g, which is known to pellet vault particles. A portion of the total cellular VR can be found in this high-speed pellet (P100), and further fractionation of the pellet over a sucrose equilibrium gradient results in cofractionation of VR and MVP, indicating that the vast majority of P100 VR is vault-associated (6). VR is also found in the non-vault-associated 100000g supernatant (S100), partly in association with the La RNA binding protein, and possibly bound to non-vault-associated TEP1 (33). To determine the accessibility of VR in unpurified vaults, we again used micrococcal nuclease digestion. In contrast to purified vaults, VR in both P100 (RNP) and deproteinized P100 (RNA) fractions was readily digested with micrococcal nuclease (Figure 1B). This suggests that the conformation of the vaults within the cell may be more open or dynamic than that of purified rat liver vaults, which are resistant to micrococcal nuclease treatment. This result, combined with the observation that VR is eventually digested by a combination of RNases even in purified vaults, prompted us to further examine the structural properties of the MVP vault shell.
VPARP and TEP1 Associate with Preformed MVP Recombinant Vaults. Vaults are predominantly cytoplasmic; however, the cellular compartment in which MVP, VPARP, TEP1, and VR assemble to form the complete vault particle is unclear, and it is also unclear if the MVP vault shell is formed prior to its association with the other vault components. As the vault-associated components (VPARP, TEP1, and VR) are also found in non-vault-associated fractions, it seems reasonable to propose that these components may shuttle into and out of the vault shell (6, 13, 14, 21). To determine whether this hypothesis is feasible given the closed capsulike structure of the vault, we turned to the Sf9 insect cell expression system where we have shown that MVP alone is sufficient for formation of recombinant vaults (15). The MVP recombinant vaults assembled via baculovirus expression appear as structurally regular as vaults purified from rat liver. Using cryoEM, it has been previously shown that capsular MVP recombinant vaults assembled via baculovirus expression is sufficient for formation of recombinant vaults (24). When VPARP and MVP lysates were mixed, VPARP in the P100 fraction shifted to the P100 pellet (Figure 2A, top right) and was copurified with MVP over a sucrose gradient (Figure 2A, top and bottom right), indicating association of VPARP with the vault particle. Identical results were obtained for TEP1-MVP lysate mixing experiments (Figure 2B). Importantly, lysate containing a TEP1 truncation that does not interact with MVP was not found in the sucrose gradient after mixing with MVP lysate (20), demonstrating that nonspecific trapping of proteins in the vault interior does not occur. Thus, the fully formed MVP shell of recombinant vaults allows for the subsequent incorporation of VPARP or TEP1.

VPARP Assembles into the Interior of the Entire Population of Vaults. Although the vast majority of the recombinant vaults purified after mixing MVP lysates with VPARP or TEP1 lysates appeared to be morphologically normal as determined by electron microscopy (see Figure 3B), it was possible that all of the VPARP and TEP1 could have assembled into a small subset of vaults that were still forming from precursors in cell lysates or from particles that were damaged during purification or incorrectly formed to some degree. To rule out these possibilities, we first looked at the amounts of VPARP that could be assembled into vaults by Coomassie staining of vaults purified from mixed lysates. The maximum number of VPARP molecules per vault in endogenous mammalian vaults is not known, nor, as previously mentioned, is it clear whether all mammalian vaults contain an equal complement of VPARP molecules. How-

FIGURE 1: Micrococcal nuclease digestion of vault RNA. (A) Purified rat liver vaults (RNP) or RNA extracted from vaults was treated with 2000 units/mL micrococcal nuclease for 0 (no enzyme), 15, 30, or 60 min. The reaction was terminated by the addition of EGTA. Control digests were incubated with micrococcal nuclease for 60 min in the presence of EGTA. (B) P100 (RNP) or RNA extracted from P100 was treated with 2000 units/mL micrococcal nuclease for 20 min (lanes 2, 3, 5, and 6) or without enzyme (lanes 1 and 4), and digestion was terminated by the addition of EGTA. Control digests included both micrococcal nuclease and EGTA (lanes 3 and 6, respectively). Reaction mixtures were extracted with phenol and chloroform, precipitated with ethanol, fractionated on a 10% acrylamide–8 M urea gel, electroblotted to Zeta GT membrane, and probed for VR. Hybridized bands were visualized by autoradiography.

FIGURE 2: VPARP and TEP1 assemble into preformed recombinant MVP-only vaults in Sf9 insect cell extracts. (A) Sf9 cell lysates containing VPARP were subjected to the vault purification protocol with (right) or without (left) mixing with extracts containing MVP recombinant vaults. A 20000g supernatant was subjected to a 100000g centrifugation, yielding a high-speed supernatant (S100) and a high-speed pellet (P100). The latter was retained for further purification. As a final purification step, samples were layered over a discontinuous sucrose gradient, and an aliquot of the subsequently pelleted gradient fractions was analyzed by Western blotting. (B) A similar analysis of TEP1 fractionation in the presence (right) or absence (left) of Sf9 cell lysates containing MVP recombinant vaults.
ever, recombinant vaults purified after lysate mixing contain substantial amounts of VPARP (Figure 3A, right lane) that appear to be equal to those obtained from either co-infection of insect cell cultures or mammalian vault purifications from tissues (1, 10). Mixing of purified MVP recombinant vaults with extracts containing VPARP followed by repurification of these vaults yielded identical results (data not shown). The high-molecular mass bands seen by Coomassie staining of purified MVP-only recombinant vaults (Figure 3A, left lane) are not VPARP since they are not detected by a VPARP antibody on Western blots (data not shown). These bands are, in fact, multimers of MVP and are recognized by anti-MVP antibodies (data not shown). Purified vaults formed with or without VPARP were visualized by negative staining followed by electron microscopy to determine any differences in the interior morphology of the two species. Analysis of recombinant vaults derived from MVP–VPARP lysate mixing, MVP–VPARP co-infection, or MVP infection alone revealed that despite the inherent variability of the negative staining technique, most vaults containing VPARP significantly excluded the uranyl acetate stain from the particle interior, indicating the presence of internal mass, and were thus easily differentiated from MVP-only vaults, which exhibit prominent interior staining which is likely due to collapse of the hollow MVP-only particles onto the carbon grids (Figure 3B). No differences in staining were visible when VPARP-containing vaults derived from lysate mixing were compared with those derived from co-infection (data not shown), indicating that VPARP localization is likely the same in the two samples.
To provide additional evidence that the majority of preformed vaults were able to incorporate VPARP, we turned to the use of electrospray mobility analysis of the vault complexes. The GEMMA separates gas-phase, predominantly singly charged particles on the basis of their differential electrophoretic mobility in air. Electrospray ionization with mass spectrometry has demonstrated the ability to assess noncovalently bound protein complexes (34, 35). With the GEMMA, ESI generates gas-phase ions representing the intact vault complexes from solution, and the ions are separated on the basis of their electrophoretic mobilities and counted by the CPC (see Experimental Procedures). On the basis of their measured mobilities, the electrophoretic mobility diameter and a molecular mass are calculated (28, 29). Although the electrophoretic mobility of a particle is governed by the size and shape of the particle, this method has also been used to characterize proteins and noncovalent protein complexes, showing a correlation between protein mass and the experimentally derived electrophoretic mobility diameter (28–30). This is based on a simple model relating molecular mass to the diameter of a sphere with an effective density (29). We reasoned that if exogenous VPARP was incorporated into only a subset of vaults, two peaks of apparent mass would be measured: one representing MVP-only vaults and one representing MVP–VPARP vaults. Analysis of vaults purified from MVP extracts with and without VPARP extract added (Figure 3C) indicates that most if not all of the vaults were able to incorporate VPARP, although the difference in apparent mass is too small to completely resolve the two peaks. The mass peak maxima were calculated from the distribution of electrophoretic mobilities corrected for the expected charge distribution of vault particles. The estimated number of VPARP copies incorporated into the vaults was calculated from replicate measurements. Figure 3C shows overlaid spectra from one set of experiments. On the basis of the estimation that there are 96 copies of CP-MVP (molecular mass of 97 kDa) per vault and the GEMMA measurement, the molecular mass of MVP-only vaults is approximately 9.3 ± 0.1 MDa (n = 3). Using the same density applied to the MVP-only vault, the molecular mass of MVP–VPARP vaults from lysate mixing and MVP–VPARP vaults from co-infection is approximately 11.2 ± 0.3 MDa (n = 3) and an average of 11.0 MDa, respectively. This corresponds to a calculated average of 9.5 VPARP copies incorporated per MVP–VPARP vault from lysate mixing and 8.3 copies of VPARP per MVP–VPARP vault from co-infection. The molecular mass distribution of both the MVP–VPARP vaults from lysate mixing and those derived from co-infection show signs of species with lower molecular masses. This indicates that there is a small fraction of these MVP–VPARP vaults that contained less than saturating levels of VPARP, particularly in MVP–VPARP vaults derived from co-infection. These results suggest that the entire population of vaults is competent to incorporate VPARP after vault shell formation.

To confirm that vaults resulting from the assembly of vault-interacting proteins into MVP recombinant vaults by mixing (in vitro) are similar to those obtained by co-infection (in vivo), we carried out fluorescence quenching experiments using GL-INT as the vault-interacting protein. GL-INT-containing recombinant vaults were purified from mixed Sf9 cell lysates containing MVP only and GL-INT only, and Sf9 cells co-infected with MVP and GL-INT encoding baculoviruses (see Experimental Procedures). Fluorescence quenching of GL upon addition of 0.5 M KCl (final concentration) has previously been shown to be substantially delayed in GL-INT vaults derived from co-infection (24). We observed similar quenching curves for GL-INT vaults derived from mixed lysates [Figure 3D (A)] and those derived from co-infected cultures [Figure 3D (B)]. In contrast, GL-INT alone in lysates was quenched immediately [Figure 3D (C)]. The delay in quenching suggests that the vault shell acts as a barrier to protect the proteins assembled into the particle interior. These data indicate that there is no difference between vaults purified from co-infected versus mixed lysates and that the incorporated proteins are inside of the vault particles.

**Cell Extracts Are Not Required for M-INT to Interact with MVP Recombinant Vaults.** We next examined whether other factors or components found in Sf9 cell extracts are required for the incorporation of exogenous proteins into the vault shell interior. For these experiments, we decided to use the C-terminal region of VPARP (amino acids 1563–1724), which is the smallest domain identified to date, that interacts with MVP, termed M-INT (for minimal size interaction domain), as it could be easily expressed and purified from *E. coli* (Figure 4A) (13, 36). Addition of a hexahistidine tag to the M-INT construct allowed for purification of the expressed 22 kDa protein using Ni–NTA chromatography.

![Figure 4: Minimal MVP interaction domain of VPARP (M-INT) that does not require Sf9 cell extracts to bind to vaults. (A) Coomassie-stained gel of *E. coli* lysates: uninduced and induced (lanes 1 and 2), column flow-through (lane 3), column wash (lane 4), and final elution of His-T7 epitope-tagged M-INT protein off the Ni–NTA resin (lane 5), which was then dialyzed into storage buffer. (B) Western blot analysis of S100, P100, and sucrose gradient fractions (described in the legend of Figure 2) after mixing of purified M-INT into Sf9 cell extracts (left) or extracts containing MVP recombinant vaults (right). (C) Mixing of 20, 40, or 100 ng of purified M-INT without (+) and with (+) 1 μg of vaults followed by high-speed centrifugation (S, high-speed supernatant; P, high-speed pellet). Pellets were resuspended in SDS sample buffer and fractionated by SDS–PAGE followed by Western blot analysis.](image-url)
Two micrograms of the highly purified protein was added to S9 cell lysate from either uninfected cells or cells infected with MVP baculovirus. M-INT fractionated in both the S100 and P100 fractions when added to extracts from uninfected cells (Figure 4B, left, S100 and P100 lanes) but did not enter the sucrose gradient (Figure 4B, left, 20–60% sucrose fractions). In contrast, when M-INT was added to vault-containing extracts, it fractionated entirely in the P100 fraction (Figure 4B, top right, S100 and P100 lanes) and copurified with MVP over a sucrose gradient (Figure 4B, top and bottom right, 20–60% sucrose fractions). We then determined whether extracts were strictly required for M-INT to assemble into vaults by incubating purified M-INT with purified MVP recombinant vaults followed by pelleting at 100000g. In the presence of MVP recombinant vaults, most of the M-INT shifted into the P100 fraction (Figure 4C, lanes 2, 6, and 10), while in the absence of vaults, M-INT remained in the supernatant (Figure 4C, lanes 3, 7, and 11). Adding increasing amounts of M-INT to 1 μg of purified MVP recombinant vaults yielded an increasing amount of protein in the pellet (Figure 4C, lanes 1, 2, 5, 6, 9, and 10). The amount of M-INT in the supernatant also increased, and this is most likely due to saturation of the VPARP (M-INT) binding sites in the MVP recombinant vaults. In summary, it appears that the interior surface of the vault barrel is accessible after a 1 h incubation, even in the absence of cell extracts, for incorporation of this 22 kDa protein.

**Thrombin Cleavage of Vault-Associated M-INT Supports Particle Conformational Dynamics.** Our data show that M-INT was readily packaged into vault particle shells. However, the possibility that the protein was bound to the outside of the particle still existed. The restricted access of the small contact quenching reagent, KCl, to the particle interior (Figure 3D) led us to conclude that GL-INT was inside the vault shell. This together with the fact that luciferase-INT was localized to the interior of the vault barrel by cryoEM reconstruction (24) strongly suggests that M-INT is also on the inside of the vault particle. As M-INT contains a thrombin cleavage site derived from the pET28 bacterial expression vector, we were able to examine the accessibility of vault-associated M-INT to thrombin, a 36 kDa protein. This approach allowed us to probe the ability of larger substances lacking a vault targeting sequence to gain access to the vault interior. Thrombin treatment of M-INT removes the pET28 vector-derived hexahistidine tag of the protein (17 amino acid residues). The cleaved and uncleaved proteins can be resolved on a 12% SDS–PAGE, transferred onto nitrocellulose, and immunoblotted for MVP (top) and the M-INT (bottom). Immunoblotting for M-INT shows both the intact and the cleaved protein. (B) Vaults treated with thrombin–agarose beads and analyzed as described above.

**DISCUSSION**

This study demonstrates that the completely formed MVP shell of recombinant vaults allows the incorporation of the very large VPARP and TEP1 proteins into the vault interior. This result was somewhat unexpected given the nature of the three-dimensional reconstructions of MVP recombinant vaults using cryoEM, which show a particle with a smooth outer surface that displays no large gaps. Thus far, the highest-resolution reconstruction of vaults (16 Å) is from the CP-MVP construct used to form vaults in this study (10). The reconstruction of the CP-MVP recombinant vault appeared to be virtually identical on the exterior to reconstructions of vaults isolated from rat tissue (10, 23), and spectroscopic results from this and previous studies using luciferase assays and fluorescence quenching indicate that the vault interior is only accessible over time (24). Therefore, some mechanism must allow for a significant modification of the vault structure that is not readily apparent in the static three-dimensional image that enables entry of large proteins into its interior. Endogenous vaults have been shown to open at the particle’s waist into two half-vault flowerlike structures when plated on polylysine-coated EM grids, and a small number of open vault halves are also seen in normal negative stain vault preparations from tissue and baculovirus-infected S9 cells (11, 15). In addition, when purified vaults are visualized by cryoEM after being rapidly frozen, some...
partially open vaults and half-vaults are seen (23). This suggests that although the vault structure overall is able to withstand a remarkable range of conditions in vitro (11), the interactions keeping the eight petals of one half-vault bound to the eight petals of the other half-vault may be less stable than the forces keeping a half-vault together. Thus a half-vault–vault equilibrium could allow entry of large molecules into the particle interior. An alternate explanation for the results obtained here is that single pairs of opposing petals (made up of multiple MVPs) on each half-vault could temporarily separate and then close, much as a door would swing open and shut, without disrupting the integrity of the particle (see Figure 6 for a schematic describing these two models). However, some other mechanism for the transient opening of holes in the vault shell may be occurring, perhaps between adjacent vault petals of the same half-vault. Insights into the precise mechanism of vault dynamics will have to await an atomic-resolution structure.

The slow digestion of VR from within the caps of purified rat liver vaults using micrococcal nuclease, or more rapid digestion using high concentrations of RNase A/RNase T1 mixtures, suggests that the shell of the vault particle is partially, but not fully, protective. It is also possible that in purified vaults, more so than in vaults in extracts, the vault RNA is protected from RNases that enter the MVP vault shell by close association with the TEP1 RNA binding protein or that the presence of TEP1, VPARP, and other protein or that the presence of TEP1, VPARP, and other interactions keeping the eight petals of one half-vault bound to the eight petals of the other half-vault may be less stable than the forces keeping a half-vault together. Thus a half-vault–vault equilibrium could allow entry of large molecules into the particle interior. An alternate explanation for the results obtained here is that single pairs of opposing petals (made up of multiple MVPs) on each half-vault could temporarily separate and then close, much as a door would swing open and shut, without disrupting the integrity of the particle (see Figure 6 for a schematic describing these two models). However, some other mechanism for the transient opening of holes in the vault shell may be occurring, perhaps between adjacent vault petals of the same half-vault. Insights into the precise mechanism of vault dynamics will have to await an atomic-resolution structure.

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Study of viral capsid structure lends plausibility to the hypothesis that vaults are dynamic particles, transiently opening and closing. In one example, the dynamics of flock house viral capsids was studied using matrix-assisted laser desorption ionization mass spectrometry (MALDI-MS) after a proteolysis time course (26). The results of this study determined that regions near the N- and C-termini of the capsid polypeptides were among the earliest sites cleaved, despite crystallographic data indicating that these regions are internal to the viral capsid. Proteolysis and related studies using chemical modification or neutralizing monoclonal antibodies to internal epitopes of poliovirus indicate that transient externalization of these internal epitopes occurs, with functional implications for cell attachment and RNA release (25). Interestingly, in flock house virus, it has been shown that the association of viral RNA alters the dynamics of the capsid structure and appears to enhance the structural integrity of the particle, although this is not evident with crystallography (37). Although the function of vaults is not known, the structural similarities to the viral capsid as “containers” made up of multimers of one or few proteins are striking, and the dynamics of internal versus external domains likely has functional significance. Of interest, one study that included pulse–chase experiments combined with immunoprecipitations of MVP and VPARP suggests that incorporation of new VPARP molecules into vaults occurs earlier than does incorporation of new MVP molecules, indicating that newly synthesized VPARP may associate with preexisting vaults (38). This result appears to be consistent with our own findings. Recently, a NMR solution structure of two-repeat units found in MVP (domains 3 and 4) has been determined, and M-INT was shown to bind to domain 4. Their model of the MVP repeats stacking suggests a distinct inner and outer surface in the vault particle with the putative VPARP binding surface on the inside of the vault (39). Additionally, the La RNA binding protein has been shown to bind to VR apart from the vault particle and to loosely associate with vaults during biochemical fractionation, leading us to hypothesize that there must be some form of VR cycling that deposits VR into the interior of the vault cap, which would seem to necessitate a dynamic vault shell (33).

Our results demonstrate that the vault is not a rigid, impermeable structure but rather a dynamic entity. Vaults could have a dual function: (i) to allow proteins to enter under certain conditions and (ii) to restrict access during other conditions so that the internal contents are protected or sequestered from other cellular factors with which they could otherwise interact. Although relatively small proteins apparently do not require cell extracts to enter vaults, it remains to be seen whether assembly factors may be required for larger proteins such as full-length VPARP and TEP1 to enter the particle interior. It will be interesting to study the factors that control opening and closing of vaults or “breathing” of the vault structure and to determine whether breathing is a important factor, not only in vault assembly but also in vault function. Vaults have been suggested to be carriers or
transporters in the cell due to both their highly conserved capsule shape and their localization at such diverse regions, including the nuclear pore complex, the leading edges of lamellipodia, adhesion plaques, the tips of neurites, and along microtubules (7, 40, 41). They have also been proposed to function as assembly scaffolds for signaling complexes (8, 9, 42). Our study demonstrates that recombiant vaults are dynamic particles and that the vault protein shell formed by MVP can transiently open to allow incorporation of other vault components and, potentially, other vault-interacting proteins. These data suggest a mechanism whereby endogenous vaults can interact with the cellular milieu.

ACKNOWLEDGMENT

We thank Stanley L. Kaufman (TSI, Inc.) for advice on the interpretation of the GEMMA data and Phoebe Stewart (Vanderbilt University, Nashville, TN) for helpful comments and critical reading of the manuscript.

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The Vault Exterior Shell Is Dynamic


BI0610552