The Role of Chaperone-subunit Usher Domain Interactions in the Mechanism of Bacterial Pilus Biogenesis Revealed by ESI-MS*

Bethny Morrissey‡, Aneika C. Leney‡, Ana Toste Rêgo§, Gilles Phan§, William J. Allen§, Denis Verger§, Gabriel Waksman§, Alison E. Ashcroft‡¶, and Sheena E. Radford‡¶

The PapC usher is a β-barrel outer membrane protein essential for assembly and secretion of P pili that are required for adhesion of pathogenic E. coli, which cause the development of pyelonephritis. Multiple protein subunits form the P pilus, the highly specific assembly of which is coordinated by the usher. Despite a wealth of structural knowledge, how the usher catalyzes subunit polymerization and orchestrates a correct and functional order of subunit assembly remain unclear. Here, the ability of the soluble N-terminal (UsherN), C-terminal (UsherC2), and Plug (UsherP) domains of the usher to bind different chaperone-subunit (PapDPapX) complexes is investigated using noncovalent electrospray ionization mass spectrometry. The results reveal that each usher domain is able to bind all six PapDPapX complexes, consistent with an active role of all three usher domains in pilus biogenesis. Using collision induced dissociation, combined with competition binding experiments and dissection of the adhesin subunit, PapG, into separate pilin and adhesin domains, the results reveal why PapG has a uniquely high affinity for the usher, which is consistent with this subunit always being displayed at the pilus tip. In addition, we show how the different soluble usher domains cooperate to coordinate and control efficient pilus assembly at the usher platform. As well as providing new information about the protein-protein interactions that determine pilus biogenesis, the results highlight the power of noncovalent MS to interrogate biological mechanisms, especially in complex mixtures of species.  *Molecular & Cellular Proteomics 11: 10.1074/mcp.M111.015289, 1–10, 2012.

Gram-negative bacteria use pili, long non-covalently assembled protein polymers, as surface-exposed appendages to mediate attachment to host cells, thereby initiating infection (1). Pili assemble at the outer membrane of the host organisms via the chaperone-usher pathway (2). Two of the most comprehensively characterized pilus systems of this pathway include type 1 pili, which attach to receptors in the bladder eventually causing cystitis (3, 4), and P pili, which attach to Galα(1–4)Gal moieties in kidney cells causing inflammation and, in some cases, urinary tract infections (5). The chromosomal pyelonephritis-associated pilus (pap) gene cluster encodes six subunits (PapG, PapF, PapE, PapK, PapA, and PapH) that assemble to form P pili (Fig. 1A). The flexible fibrillum contains at its tip only one copy of the PapG subunit which, uniquely, contains both an adhesin and a pilin domain (Fig. 1A). This subunit is required to be displayed at the pilus tip for host adhesion (6). PapG is joined to one copy of PapF, which is then linked to 5–10 copies of PapE to form the flexible pilus tip. A single PapK subunit joins the flexible tip to the second segment, a right-handed, rigid, helical rod, consisting of thousands of PapA subunits (Fig. 1A). Pilus assembly is arrested by the incorporation of one copy of the PapH subunit which prevents further assembly and locks the fully assembled pili into the usher complex in the outer membrane (7–10).

Assembly of the different subunits in a precise order is required to form functional pili. Pilus biogenesis, thus, is a highly organized process which is mediated by a large β-barrel transmembrane protein known as the usher (PapC) (Fig. 1B). Subunits (PapX) enter the periplasm via the secYEG general secretory pathway (11) whereupon they each bind to a molecule of the same chaperone, PapD, forming a PapDPapX chaperone-subunit complex (supplemental Fig. S1A, S1C). Chaperone-subunit binding is required to prevent the subunit domain from misfolding, premature polymerization, and degradation (9, 12). Each PapDPapX complex is formed by the chaperone donating one of its β-strands (Gβ) to the pilin subunit, to complete its C-terminally truncated immunoglobulin (Ig) fold, through a process known as donor strand complementation (13) (supplemental Fig. S1A).

To bring about pilus assembly, PapDPapX complexes are targeted to the usher at the bacterial outer membrane (Fig. 1B). The usher is an 809 residue protein comprised of a
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24-stranded β-barrel pore that, in the inactive state, is blocked by its plug domain (UsherP) (magenta domain Fig. 1C, left) (14–15). Chaperone-subunits are first bound by the soluble N-terminal domain of the usher (UsherN) (red domain Fig. 1C) (16). Binding of PapDPapG (the first complex recruited at the usher) to UsherN activates the usher and causes release of UsherP into the periplasm, opening the translocation pore (Fig. 1B, right) (14, 15). Subunits are released from the chaperone when the N-terminal extension (Nte) of the subunit next in assembly displaces the donated chaperone G1, β-strand in a process termed donor strand exchange (DSE) (9) (supplemental Fig. S1B). During, or after, DSE subunits are released from UsherN and passed (directly or indirectly) to a second pair of soluble domains located toward the C-terminal region of the usher protein (UsherC1 and UsherC2) (green domains Fig. 1B, right) (15), whereupon they emerge through the pore of the usher (15) creating the new, growing pilus. This process continues, (catalyzed by the usher) (17), until a PapDPapH complex reaches the usher and undergoes DSE, terminating pilus growth (9).

The relative rates of DSE between all 30 possible pairs of P pilus subunits and their Ntes have been investigated using electrospray ionization mass spectrometry (ESI-MS) (18, 19). These experiments revealed that cognate subunit-Nte pairs undergo more rapid rates of DSE than their noncognate counterparts, suggesting that the relative rates of DSE contribute toward subunit ordering in pilus biogenesis. However, the extent of subunit discrimination and the rates of DSE observed in vitro are too small or too slow, respectively, to explain the specificity and rate of pilus biogenesis in vivo (20–21). This is consistent with a role of the usher protein in catalysis of pilus biogenesis (17) and in subunit discrimination (22, 23). How this is accomplished, however, remains unclear, despite recent crystal structures of the usher complex both with and without chaperone-subunits bound (14, 15, 22–25) (Fig. 1B).

Here we describe the use of noncovalent ESI-MS to monitor interactions of the three soluble usher domains (UsherN, UsherC2, and UsherP) (Fig. 1C) individually with all six PapDPapX complexes (supplemental Fig. S1C) (UsherC1 of PapC could not be produced by recombinant expression). Combined with collision induced dissociation tandem mass spectrometry (CID-MS/MS) to assess the relative gas phase stability of the different complexes formed, and competition binding experiments, also monitored by ESI-MS, the results reveal the ability of all three soluble usher domains to bind all six chaperone-subunit complexes, suggesting their involvement in the movement of the chaperone-subunit complexes around the usher to bring about the correct ordering of subunits into the usher pore during pilus biogenesis.

**EXPERIMENTAL PROCEDURES**

Expression and purification of soluble usher domains and PapDPapX complexes—PapDPapG complexes—PapDPapG total intensity = PapDPapG (18, 26, 27), PapGad (28), PapDPapGcarb (18), PapDPapGad (9), PapDPapG (9), PapDPapGHis1 (10) and PapDPapGHis1 (10), and the UsherC2 domain (24) were cloned, expressed and purified as previously reported. Production of PapDPapGHis1, UsherN, UsherN24–131, and UsherP was described in the Supplementary Information.

Formation of Ternary PapDPapUXusherY Complexes—PapDPapX complexes and each soluble usher domain were dialyzed separately overnight into 5 mM ammonium acetate (pH 6.8) after which the usher domain (final concentration of 20 μM) (in the same buffer) was added to each chaperone-subunit complex (final concentration 10 μM) to a total volume of 50 μl in 5 mM ammonium acetate (pH 6.8). After 30 min at ambient temperature, 6 μl of sample was removed and analyzed by ESI-MS. The concentration and pH of ammonium acetate were selected based on previous optimization studies for buffer conditions for analysis of the chaperone-subunit complexes (18).

**Competition Experiments**—Competition experiments in which the UsherN domain was added to a mixture of PapDPapG and PapDPapA involved dialyzing all components overnight into 5 mM ammonium acetate at pH 6.8, followed by the analysis of the different protein complexes after mixing on the subsequent day. All of these experiments were performed in series on the same day. Increasing concentrations of UsherN (0–10 μM) were added to equimolar amounts (5 μM) of PapDPapG and PapDPapA. After 15 min, an aliquot was removed and analyzed by ESI-MS. Comparison of the relative ion intensity of different complexes was calculated as follows: relative ion intensity of PapDPapG-UsherN = PapDPapG(UsherN total intensity)/PapDPapG total intensity + PapDPapG (UsherN total intensity). Likewise, relative intensities of PapDPapA-UsherN were calculated by the following equation: relative ion intensity of PapDPapA-UsherN = PapDPapA (UsherN total intensity)/PapDPapA total intensity + PapDPapA (UsherN total intensity).

For competition experiments involving two usher domains and one chaperone-subunit complex, after overnight dialysis the first usher domain (10 μM) was added to each chaperone-subunit complex (10 μM) in a total volume of 20 μl in 5 mM ammonium acetate (pH 6.8). After 2 h at ambient temperature, 6 μl of sample was removed and analyzed by ESI-MS. The second usher domain or control protein (10 μM) was then added and after a further 2 h at ambient temperature, 6 μl of sample was removed and analyzed by ESI-MS.

**Mass Spectrometry Data Acquisition and Interpretation**—Ternary (chaperone-subunit-usher(N, C2, or P)) and competitive (PapDPapG + UsherN, or chaperone-subunit-usher(N, C2, or P)) complexes were identified using a LTQ Orbitrap Premier (Micromass UK Ltd., Waters Corporation, Manchester, UK) time-of-flight mass spectrometer. ESI-CID-MS/MS experiments were performed with a Synapt HDMS, hybrid quadrupole-ims-tof mass spectrometer (Micromass UK Ltd., Waters Corporation, Manchester, UK). Both mass spectrometers are interfaced with a NanoMate (Advion Biosystems Inc., Ithaca, NY) nanoESI autosampling device for sample introduction and ionization. Positive nanoESI with a capillary voltage of 1.75 kV and a nitrogen nebulising gas pressure of 0.5 p.s.i. was used. A cone voltage of 70 V with a source temperature of 50 °C was applied. For identification of ternary and quaternary complexes, spectra were acquired over 1 min with the analyzer set to transmit m/z 1000–5000 for all complexes except those containing the PapDPapG subunit, in which case ions were acquired over the range of m/z 1000–7000. MassLynx v.4.1 software was used to process raw data.

1 The abbreviations used are: UsherP, plug usher domain; ESI-MS, electrospray ionization mass spectrometry; CID, collision induced dissociation; MS/MS, tandem mass spectrometry; Pap, pyelonephritis associated pilus; DSE, donor strand exchange; Nte, N-terminal extension; PapDPapX, chaperone-subunitX; where X = PapG, PapF, PapK, PapE, PapA, or PapH; UsherN, N-terminal usher domain; UsherC1/2, C-terminal usher domains.
ESI-CID-MS/MS experiments were performed by selecting and isolating ions representing the ternary complex in the trap collision cell that was filled with argon at a pressure of 1.18 \times 10^{-2} mbar. Increasing collision energy was applied to the trap collision cell in 10 eV increments until the product ions were completely dissociated into the fragment ions. Profiles describing the fragmentation pathway were generated. These display the fraction of ternary ion intensity (as a percentage of total ion intensity) versus normalized collision energy 

\[ NCE = \frac{E \times A_r \times z}{A_r + C_m} \]

where \( E \) is the collision energy, \( A_r \) is the mass of Argon, \( C_m \) is the mass of the complex and \( z \) the charge of the precursor ions.

**RESULTS**

**Chaperone-Subunit-Usher Ternary Complexes Identified by ESI-MS**—Structural analyses of the homologous usher proteins, PapC and FimD (from P pilus and type I pilus, respectively) have revealed that these proteins contain five functional domains: a membrane-embedded translocation domain and four soluble domains that reside in the periplasm in the activated state (Figs. 1B, C) (14, 15). The latter include the 131-residue N-terminal domain (UsherN), two C-terminal domains (UsherC1, 84 residues; UsherC2, 85 residues) and the 83-residue plug domain (UsherP) (Fig. 1C) (15). To determine whether the soluble Pap usher domains are able to bind different PapDPapX complexes, an ESI-MS in vitro assay was developed to detect interactions between the different soluble usher domains and different chaperone-subunit pairs. All six PapDPapX complexes (supplemental Fig. S1C) and three soluble usher domains (UsherN, UsherP, and UsherC2; Fig. 1C) were expressed and purified as previously reported (18) and their molecular masses confirmed by ESI-MS (supplemental Text and Fig. S2). Efforts to produce UsherC1 were not successful, presumably because this domain makes intimate interactions with the lumen of the usher translocation domain (Fig. 1B) and is therefore unstable in isolation. Each of the soluble usher domains was mixed separately with each PapDPapX complex and binding was monitored using noncovalent ESI-MS under ambient conditions (Experimental Procedures). Strikingly, ternary complexes of all six PapDPapX complexes with UsherN, UsherC2, or UsherP were identified using ESI-MS, resulting in ions indicative of complexes representing one usher domain bound to one chaperone-subunit in all cases (red peaks, Fig. 2, and supplemental Fig. S3). The ion series representing the ternary complex (red peaks, Fig. 2, and supplemental Fig. S3) appear interspersed with the ion series representing the free chaperone-subunit complex (cyan peaks, Fig. 2 and supplemental Fig. S3). The measured molecular masses of the ternary complex were in close agreement (within 0.03%) with the expected masses (supplemental Table S1).

Despite clear evidence of binding between all of the PapDPapX complexes with each usher domain, different ion intensities are observed for the different complexes formed (supplemental Fig. S3). This likely reflects differences in binding affinity, complex stability and/or ionization efficiency. In accord with the latter proposition, control experiments showed that the three usher domains give very different MS responses; for example, the UsherC domain seemingly generates a lower ion count than either the UsherN or the UsherP domain when pairs of these proteins are mixed at equimolar concentrations in the absence of PapDPapX complexes (supplemental Fig. S4). Also, the var-

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**Fig. 1. Schematic of subunits comprising a P pilus and structure of the usher platform.** A, Organization of the P pilus, which includes one copy each of PapG (red), PapF (orange), PapK (green), and PapH (dark blue), five to ten copies of the PapE (yellow), and up to one thousand copies of PapA (cyan). The chaperone, PapD (D, brown), usher (C, pink), periplasm (P), extracellular space (Ex), and bacterial outer membrane (OM) are indicated. The PapG adhesin (left, pdb 1J8S (28)) and pilin (right, pdb 3MEO) domains are shown. B, Structure of the usher FimD protein in the apo (left, pdb 3OHN (15)) and activated (right, pdb 3RFZ (15)) forms. The soluble N-terminal (red), plug (magenta), and C-terminal (C1; lime, C2; dark green) domains are shown. Note that the plug domain is released from the usher lumen into the periplasm in the activated form. The structure of the activated form also contains a chaperone-subunit (brown-blue) complex bound to the C-terminal usher domains. C, Domain organization of the PapC usher colored as in (B).
ious chaperone-subunits (in the absence of usher) do not ionize with identical efficiencies (supplemental Fig. S5). It is not unexpected, therefore, that the ternary complexes arising from different usher domains and different chaperone-subunit-usher domain complexes give rise to differing MS responses that will not reflect their relative abundances in solution quantitatively, and hence nor their $K_D$ values.

Previous reports have shown direct interactions of the chaperone-adhesin complex (PapDPapG) with the C-terminal domain (UsherC2) of the usher (15, 29), but no evidence has been reported for interactions of other chaperone-subunit complexes with this domain, nor for any PapDPapX complex with UsherP. The interactions observed here by ESI-MS appear to be specific, since there was no evidence for complexes with a molar ratio of >1 usher domain binding to each chaperone-subunit complex even when the usher domain is added in excess (supplemental Figs. S4 and S5). In the crystal structure of UsherN bound to a chaperone-pilin complex, the N-terminal residues 2–23 of UsherN form a flexible tail that contacts the chaperone (22). Consistent with this, residues 2–11 of UsherN have been shown to be required for high affinity binding to chaperone-subunit complexes (16). To determine whether this is also the case in the experiments presented here, the ability of a truncated UsherN construct, UsherN$^{24-131}$ (supplemental Text) to bind to all six PapDPapX complexes was also investigated. The ESI-MS results showed that UsherN$^{24-131}$ is still able to form ternary complexes with all six PapDPapX complexes, indicating that the presence of residues 1–23 is not obligate for chaperone-subunit binding (red peaks, supplemental Fig. S7A). In addition to the known ability of UsherN to bind all six PapDPapX pairs (16, 22, 30), the results presented here reveal that UsherC2 and UsherP also recognize all six different chaperone-subunit complexes, suggesting that each of the soluble usher domains may play a role in the assembly of type P pili.

**Differences in the Relative Stabilities of Chaperone-Usher-Subunit Complexes**—Because all soluble usher domains were found to be capable of forming stable interactions with each PapDPapX complex (red peaks, Fig. 2, and supplemental Fig. S3), the difference in the gas phase stability of the different PapDPapX-UsherX ternary complexes was next determined using CID-MS/MS. These experiments are indicative of the relative gas phase stabilities of the different complexes formed (31). CID-MS/MS has been used widely to compare gas phase stabilities of protein complexes (32–35), including the stability effect of anions (34), mutations (36), and ligand binding (37–38) on protein complexes. In this study, this ubiquitous technique has been used to compare the behavior of differing chaperone-subunits complex bound to an usher domain in a qualitative context. In this experiment, ions representing the ternary complex were selected as the precursor ions. Accordingly, the collision energy applied to effect dissociation of different precursor ions was increased until complete dissociation of each ternary complex was observed. The fraction of the precursor ion intensity remaining compared with the total ion intensity was then plotted as a function of normalized collision energy (Fig. 3). Consistent with the known high affinity of PapDPapG for UsherN determined using Surface Plasmon Resonance (SPR) experiments (39), PapDPapGUsherN was also found to be the most stable ternary complex in vacuo (Fig. 3A), with the other five PapDPapX-UsherN complexes showing substantially weaker, but nonetheless significant, gas phase stability relative to PapDPapG-UsherN (Fig. 3A, inset).

The ion intensities of the ternary complexes may not reflect in-solution abundances (31). To determine whether the relative gas phase stability is reflective of solution phase binding affinity for the noncovalent complexes investigated here, the
interactions of PapDPapG and PapDPapA with UsherN were followed in more detail using ESI-MS. These two chaperone-subunit complexes were chosen specifically for this analysis since the binding of these complexes to UsherN has been determined previously using SPR (39). These solution phase binding data revealed that UsherN binds significantly more tightly to PapDPapG compared with PapDPapA (K_d values of 9.1 × 10^{-9} M and 1.8 × 10^{-7} M, respectively (39)), corroborating the CID-MS/MS data which show an enhanced gas phase stability for the PapDPapGUsherN complex compared with PapDPapAUsherN (Fig 3A). To demonstrate the different binding affinities of these complexes further using ESI-MS, PapDPapG and PapDPapA were mixed in an equimolar ratio and their relative ion abundances were determined (Fig. 4, top panel). ESI-MS analysis revealed that PapDPapG ionizes more efficiently than PapA, as previously discussed. This equimolar mixture of PapDPapG and PapDPapA was then mixed with UsherN either in a 1:1:1 or 1:1:2 molar ratio of PapDPapG, PapDPapA, and UsherN, respectively, and the relative ion intensities of the different ternary complexes were measured and compared (Fig. 4, lower panels). The results revealed that when Usher N is added to PapDPapG and PapDPapA in a 1:1:1 ratio, UsherN is found predominantly bound to PapDPapG whereas very little, if any, binding was observed to PapDPapA (Fig. 4, central panel). By contrast, when UsherN was added in a 1:1 ratio to either PapDPapG or PapDPapA (without the competing chaperone-subunit complex), significant ion intensity corresponding to each ternary complex was observed (Figs. 4 and supplemental Fig. S3). Only when an excess of UsherN is added to PapDPapG/PapDPapA in a molar ratio of 1:1.2 is an observable ion intensity of the ternary PapDPapAUsherN complex detected (Figs. 4 and supplemental Fig. S5), thus reinforcing the ability of ESI-MS to reveal the relative binding affinity of different

**Fig. 3.** Stability of different PapDPapXUsherX complexes determined using CID-MS/MS. A, CID-MS/MS fragmentation profiles of ternary complexes involving all six PapDPapX complexes bound to UsherN. B, Fragmentation profiles of PapDPapGUsherN (square), PapDPapGUsherN_{24–131} (triangle), PapDPapG_UsherN (diamond), and PapG_p_UsherN (circle). C, Fragmentation profiles of different PapDPapX complexes bound to UsherP and (D) but bound to UsherC2. Inset: bar graphs representing the normalized collision energy required to fragment each ternary ion to 50% of its original intensity. Error bars represent the S.E. of four replicate measurements.
complexes, provided that suitable care is taken to measure the relative ionization efficiency of the different species formed, as well as to compare different species within carefully controlled experimental parameters. The data are thus consistent with the SPR and CID-MS/MS data, all of which indicate tighter binding of UsherN for PapDPapG.

To determine the contribution of the unstructured N-terminal region of UsherN to the stability of the complexes formed between this usher domain and PapDPapG, ternary complexes were formed between PapDPapG and the truncated UsherN24–131 construct, and the gas phase stability of the resulting complex was determined using CID-MS/MS (Fig. 3B). Consistent with the known role of the N-terminal region of UsherN in determining high affinity PapDPapG binding (16), the gas phase stability of the PapDPapGUsN complex was reduced substantially when the unstructured N-terminal region was deleted (Fig. 3B). Interestingly, the stability of the other five chaperone-subunit complexes was not affected by the absence of the residues 1–23 in UsherN (supplemental Fig. S7A, S7B).

Although five of the six PapX subunits contain only a single pilin domain, PapG is unique among the pilin subunits in that it contains two domains, a pilin domain (PapGp) and an adhesin domain (PapGad) (Fig. 1, supplemental Fig. S1C). Thus, we also tested complex formation and stability to gas phase dissociation of a PapDPapGpUsherN complex lacking the adhesin domain of PapG (Fig. 3B triangles). Remarkably, complex stability is reduced substantially in this construct relative to its intact PapDPapG counterpart (Fig. 3B squares). These results reveal a hitherto unsuspected role of the PapG adhesin domain in enhancing the binding affinity of PapG for UsherN. Indeed, without the adhesin domain, the stability of the PapDPapGpUsherN complex is similar to that of other PapDPapX complexes (Fig. 3B). To confirm the ability of UsherN to bind to the adhesin domain alone, PapGadhesin (Ga) (residues 1–196 of PapG, Fig. 1, supplemental Fig. S1C) was constructed (Experimental Procedures) (28) and the binding of this domain to UsherN was determined by ESI-MS (supplemental Fig. S8). The stability of the binary complex of PapGad with UsherN was also measured using CID-MS/MS (Fig. 3B circles), revealing that this complex has similar gas phase stability to PapDPapGp. The data strongly suggest, therefore, that a binding interface exists between the UsherN domain of PapC and the adhesin domain of PapG. This inter-

Fig. 4. Competition binding experiments with UsherN and mixtures of PapDPapG and PapDPapA. A, ESI-MS spectra of an equimolar mixture of PapDPapG + PapDPapA (PapDPapG (cyan), PapDPapA (green)). Bar charts showing the relative ion intensities of each complex within the mixture are shown alongside. B, ESI-MS spectra of equimolar PapDPapG + PapDPapA to which UsherN in a 1:1:1 molar ratio is added. Ions arising from the different complexes are colored: PapDPapG (cyan), PapDPapA (green), PapDPapAUsherN (purple), PapDPapGUsN (red). Bar chart showing the relative ion intensities of ternary complexes PapDPapGUsN (red) in samples that either lack (labeled PapDPapG + UN) or contain an equimolar concentration of competing PapDPapA (labeled PapDPapG + DA/UN) are shown alongside. Similarly, the ion intensity of PapDPapAUsherN (purple) without added PapDPapG (labeled PapDPapA + UN) or in the presence of the competing complex (labeled PapDPapA + DG/UN) are also shown (see Experimental Procedures). C, ESI-MS spectra of equimolar PapDPapG + PapDPapA to which UsherN in a 1:2 molar ratio is added, with relative ion intensities shown in the corresponding bar chart. The ions are colored and labeled as in (B). In all cases errors shown are the S.E. over several replicate measurements.
action rationalizes the remarkable stability of the PapDPapG
UsherN ternary complex that may facilitate activation of the
usher with the PapDPapG complex, which is required to ini-
tiate the formation of a functional pilus (40).

In parallel with measurements of the gas phase stability of
ternary complexes of different chaperone-subunits with
UsherN, similar measurements were performed for all six
PapDPapX complexes with UsherP and UsherC2. All of these
complexes were stable to gas phase dissociation, resulting in
measured stabilities similar to those formed between UsherN
and PapDPapF/E/K/A/H, but weaker than the complex
formed between UsherN and PapDPapG (compare bar
graphs in Figs. 3A, 3C, and 3D). Small, but significant, differ-
ences in the stability of the ternary complexes involving
UsherP and UsherC2 with the six PapDPapX complexes were
observed (Fig. 3C, 3D), suggesting that the binding interface
must involve at least some interactions with the subunit do-
main itself as the chaperone is the same across the PapD-
PapX complexes (supplemental Fig. S1C). Consistent with
this, dissociation of the ternary complexes yielded ions rep-
resenting the PapDPapX complex in addition to ions indicat-
ing the presence of free PapD, free PapX and free usher
domains. Ions that were consistent with both PapDUsherY
and PapUXsherY binary complexes were also detected.
These findings support the proposal of a shared binding in-
terface for all three usher domains with both the chaperone
and subunit components of the complexes, consistent with
recent crystallographic information of the equivalent subunits
from type 1 pilus (FimCFimH) binding to the N- and C-terminal
domains of the Fil usher, FimD (15, 22). During the CID-
MS/MS fragmentation of the ternary complexes, the UsherN
domain was dissociated from the PapDPapX chaperone-sub-
unit complex, following an expansion in its collision cross-
sectional area, consistent with dissociation mechanisms re-
ported elsewhere (35, 41–43).

**Competition Between Different Usher Domains for Chaper-
one-Subunit Binding**—Recent crystallographic analysis of the
FimD usher in the act of DSE and subunit translocation through
the pore of the usher has suggested that UsherN and
UsherC2 compete for binding to each chaperone-subunit
complex (15). Although UsherN has been proposed to be
essential for receiving (16, 22) and discriminating (30, 39, 44)
between the incoming chaperone-subunit complexes, usher
domains C1 and C2 have been proposed to bind to the
chaperone-subunit complex after DSE, releasing UsherN to
accept the next incoming subunit (15). How handover of the
chaperone-subunit complex between usher domains is con-
ducted, however, remains unclear. Two scenarios can be en-
visioned: direct competition of UsherN and UsherC1/2 for a
common binding site on the chaperone:subunit complex, or
transfer between UsherN and UsherC1/2 via an intermediary
domain or surface.

To determine whether UsherN, UsherP, and UsherC2 com-
pete for a common binding site on each chaperone-subunit
complex, the ability of each usher domain to bind simultane-
ously to PapDPapG or PapDPapF was investigated. Accord-
ingly, equimolar concentrations of two different usher do-
 mains were added sequentially to PapDPapG or PapDPapF
and the resulting mixtures were analyzed before, and after,
addition of the second usher domain using ESI-MS. When
equimolar amounts of UsherC2 or UsherP were added to
complexes of PapDPapG or PapDPapF containing prebound
UsherN (Fig. 5), the results revealed a significant depletion in
the fraction of PapDPapGUsherN and PapDPapFUsherN ions
compared with their unbound chaperone-subunit predeces-
sors, suggesting that these domains can displace UsherN via
direct competition for a common binding site or an allosteric
shift. Importantly, when control proteins (UsherC2 of FimD or
hen egg white lysozyme) were added, the fraction of PapD-
PapGUsherN to unbound PapDPapG was not depleted,
suggesting the competition for UsherN is specific to the
PapC usher domains (supplemental Fig. S9). Quaternary
complexes were observed for PapDPapGF/UsherN+UsherC2
and PapDPapG+UsherN+UsherP, but not for PapDPapGF/UsherC2+UsherP (Fig. 5A). These potential
intermediates may be “snapshots” of a possible displace-
ment mechanism occurring between the two domains, or
evidence of a handover process between UsherN, UsherP,
and UsherC2.

**DISCUSSION**

*The Adhesin Domain is not an Innocent Bystander in Initi-
ating Pilus Formation*—The UsherN domain of PapC is known
to be the first site of interaction of the usher with PapDPapG,
the binding of which has to occur first in order to assemble a
functional pilus (15, 16). In the case of the usher in the closely
related Type 1 pilus, (supplemental Fig. S10) FimD, activation
has been shown to require binding of the FimH subunit that
contains the adhesin domain (supplemental Fig. S10), ensur-
ing that this subunit is displayed at the pilus tip as required for
functionality (45, 46). By contrast with the strict requirement of
FimD for activation with FimH, PapC is less discriminating and
may be activated by the binding of chaperone-subunit com-
plexes other than PapDPapG (16, 40). Here we show using
CID-MS/MS of the PapDPapXUsherN ternary complexes that
the PapDPapGUsherNC complex is substantially more stable to
gas phase induced dissociation than the other ternary com-
plexes (Fig. 3A), consistent with the known high affinity of this
complex determined using SPR (39, 44, 47). Dissection of
PapG into its constituent adhesin and pilin domains, together
with analysis of binding of different chaperone-subunit pairs
to UsherN and its truncated variant UsherN_{24–131}, revealed a
dramatic reduction in stability when the UsherN_{24–131} con-
struct was bound to PapDPapG (Fig. 3B), or when an adhesin-
deleted PapDPapG complex was added to UsherN. These
data indicate a hitherto unsuspected role of the adhesin
domain in contributing to the high affinity binding of PapDPapG
to UsherN. Interestingly, crystallographic analysis of the com-
plex formed between the FimD UsherN domain bound to FimCFimHp (in which the adhesin domain of FimH is deleted) revealed that the first 23 residues of UsherN interact with the chaperone and not with the pilin domain (22). This may result from the lack of the adhesin domain in this complex, or could reflect differences in the manners by which the different UsherN domains of PapC and FimD recognize their initiating chaperone-subunit targets.

**Coordinated Participation of UsherN, UsherP, and UsherC2 in Pilus Biogenesis**—The usher plays numerous key roles in pilus biogenesis, including catalysis of DSE (17) and facilitation of subunit ordering (39, 40, 44). The usher is critical for pilus construction, as without this machinery pili are no longer assembled or secreted (20, 48, 49). Here, using noncovalent ESI-MS, we show that each soluble usher domain (UsherN, UsherC2, and UsherP) is able to bind each of the six PapD-PapX complexes, suggestive of a potential role for all three soluble usher domains in pilus biogenesis (Fig. 2 and supplemental Fig. S3). To date, no interactions of UsherC or UsherP with chaperone-subunit complexes have been reported for the Pap system, highlighting the power of ESI-MS to observe these interactions. These observations are consistent with a recent crystal structure of the structurally and functionally homologous FimD usher in complex with the initiating FimCFimH chaperone-subunit complex, which reveals close contacts between FimCFimH with the UsherC1 and UsherC2 domains and less extended, but nevertheless significant, contacts with the plug domain (15). Furthermore, previous deletion experiments have shown that all three soluble usher domains are required for pilus assembly in vivo (14, 23, 29, 40, 50). Therefore, a well-orchestrated and specific series of interactions between the chaperone-subunits and all three soluble usher domains appears to be required to ensure effective pilus biogenesis.

**A Refined Model for Usher Discrimination in Pilus Biogenesis**—From the data presented, a model of the series of interactions between each PapD-PapX complex with the usher domains during pilus biogenesis can be constructed (Fig. 6). Pilus assembly commences with the PapD-PapG complex binding to the UsherN domain of PapC (Fig. 6 (step A)) (15, 16). Binding of PapD-PapG causes activation of PapC and the consequent displacement of UsherP from the lumen of the usher into the periplasm (Fig. 6 (step B)) (15). The data presented here show that UsherP and UsherC2 are each able to bind all six PapD-PapX complexes (Fig. 2, supplemental Fig. S3), indicating a potential role of these domains in pilus assembly. For UsherP this could involve facilitating the transfer of the chaperone-subunit complexes between the N- and C-terminal usher domains, consistent with the ability of this domain to bind all six chaperone-subunit complexes (supplemental Fig. S3), in addition to displacing the incoming subunit from UsherN (Fig. 5). Once the PapD-PapXUsherN ternary complex is formed, the chaperone-subunit complex could be passed directly to the C-terminal usher domains (Fig. 6, pathway I, (Step C)), or this transfer could be mediated via UsherP (Fig. 6, pathway ii). In support of the latter, previous experiments have demonstrated a requirement of the plug domain.
A consequence of UsherP domains binding simultaneously with ternary complexes, the detection of quaternary complexes involving the UsherC domain for successful pilus formation, but not for the structural mechanism (25, 51). Also, potential intermediates of these handover processes are observed by the detection of quaternary complexes involving the UsherC and UsherP domains binding simultaneously with ternary complexes involving UsherN (Fig. 5A). As a consequence of this transfer, UsherN would be released for recruitment of the next chaperone-subunit complex (Fig. 6 (step D)), whereas binding of the newly transferred chaperone-subunit complex to UsherC1/2 positions the Nte of the incoming subunit directly over the P5 pocket of PapG, allowing DSE to occur (Fig. 6 (step D)) (15).

In summary, the data presented here add to the growing insights into the molecular mechanism of pilus biogenesis captured by crystallographic snapshots of different points in the assembly mechanism (14, 15, 23). Specifically, using ESI-MS we have revealed a new and hitherto unsuspected role of the adhesin domain in enhancing the affinity of PapG for UsherN, which may play a part in ensuring that this domain is used to initiate pilus formation by activating the usher. In addition, we reveal a potential role of the plug domain in orchestrating the transfer of each chaperone-subunit complex between the UsherN and UsherC domains, ensuring efficient transfer of subunits in the assembly line, and smooth handover between the UsherN and UsherC domains. Utilizing the sensitivity and ability of ESI-MS to monitor and analyze individual species within heterogeneous assemblies and to compare the relative strength of noncovalent interactions using both CID-MS/MS and competition experiments, binding interactions between usher domains and chaperone-subunits have been investigated. By extending this approach to interrogate interactions of chaperone-subunits with the whole membrane-embedded usher protein more insights into pilus biogenesis are sure to be revealed.

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This article contains supplemental Figs. S1 to S10, Table S1, and Experimental Procedures.

To whom correspondence should be addressed: Astbury Centre for Structural Molecular Biology, Institute of Molecular and Cellular Biology, The University of Leeds, Leeds, LS2 9JT, UK. Tel.: ++44 113 343 7273; Fax ++44 113 343 7273; E-mail a.e.ashcroft@leeds.ac.uk.

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