A small surface hydrophobic stripe in the coiled-coil domain of type I keratins mediates tetramer stability

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Intermediate filaments (IFs) are fibrous polymers encoded by a large family of differentially expressed genes that provide crucial structural support in the cytoplasm and nucleus in higher eukaryotes. The mechanisms involved in bringing together elongated coiled-coil dimers to form an IF are poorly defined. Available evidence suggests that tetramer subunits play a key role during IF assembly and regulation. Through molecular modeling and site-directed mutagenesis, we document a hitherto unnoticed hydrophobic stripe exposed at the surface of coiled-coil keratin heterodimers that contributes to the extraordinary stability of heterotetramers. The inability of K16 to form urea-stable tetramers in vitro correlates with an increase in its turnover rate in vivo. The data presented support a specific conformation for the assembly competent IF tetramer, provide a molecular basis for their differential stability in vitro, and point to the physiological relevance associated with this property in vivo.

Introduction

Intermediate filaments (IFs) are flexible intracellular fibrous polymers that provide resilience to the cells in which they are expressed (Fuchs and Cleveland, 1998; Omary et al., 2004). The structural support function of IFs is made possible by their unique ability to sustain relatively large deformations without breaking (Janmey, 1991; Yamada et al., 2002). Loss of this function through inherited mutations underlies a large variety of rare diseases in which affected cells and tissues are often fragile and cannot sustain mechanical stress (Omary et al., 2004). Additionally, IFs can also modulate the response to chemical stress, pro-apoptotic, and other signals, a newly defined function that involves regulated interactions with signaling effectors (Coulombe and Wong, 2004).

The human genome contains at least 67 distinct genes encoding proteins able to self-polymerize into 10-12-nm-wide IFs that are regulated in a tissue-, cell type–, and differentiation-specific fashion (Hesse et al., 2001). All IF proteins (M, 40–240 kD) share a tripartite domain organization consisting of a central \( \alpha \)-helical “rod” flanked by nonhelical head and tail domains. In cytoplasmic IF proteins, the rod domain is 310 aa residues long and features long-range heptad repeats (abcdefg) in which amino acid residues located in the first a and fourth d positions are hydrophobic or apolar, leading to the “knob and hole” packing of two \( \alpha \)-helices into a stable coiled-coil dimer (Crick, 1952; Cohen and Parry, 1990). The heptad repeats are interrupted by short linker sequences at three conserved locations, segmenting the rod domain into coils 1A, 1B, 2A, and 2B (35, 101, 19, and 121 residues long, respectively; Parry and Steinert, 1999). The head and tail domains exhibit variable primary and secondary structure and are substrates for phosphorylation and other modifications that regulate IF polymer assembly, dynamics, and interactions with other proteins (Coulombe and Omary, 2002).

Resolving the high resolution structure of F-actin and microtubule fibrous polymers has catapulted these research fields to new heights. Such a detailed understanding is lacking for IFs, owing to the elongated shape and polymerization-prone nature of their constituent proteins, and structural polymorphism (Strelkov et al., 2001). Scanning transmission electron microscopy has shown that on average the IF polymer backbone consists of 16 coiled-coil dimers in cross section (Herrmann et al., 1999). Dimers are structurally elongated (46-nm length; 2–3-nm width; Quinlan et al., 1984) and polar, because of the parallel and in-register alignment of \( \alpha \)-helices. Packing 16 dimers into a smooth surfaced, 10–12-nm filament represents a tour de force that remains poorly understood. For cytoplasmic IFs, dimers interact along their lateral surfaces with an antiparallel orientation to form apolar tetramers. In vitro, further lateral interactions between tetramers yield unit length filaments (ULFs, 16-nm width; 60-nm length), which then anneal and compact to give rise to mature IFs (Strelkov et al., 2003). Beyond dimer formation, the interactions presiding over the polymerization of IF proteins are less understood.
Results

Like its human orthologue, mouse K16 forms unstable tetramers with type II partner K6

We compared the ability of mK16 to form stable heterotetramers in vitro to that of hK16 and related type I keratins, hK14 and mK17. Purified types I and II keratins were mixed in the presence of 6.5 M urea and low salt and subjected to an anion exchange chromatography assay that resolves monomers, heterodimers, and heterotetramers (Wawersik et al., 1997). Whereas hK14/mK6 and mK17/mK6 elute largely as heterotetramers, both hK16/mK6 and mK16/mK6 elute primarily as monomers and heterodimers (Fig. 1 a). Complexes containing mouse or human K6 paralogues and K17 display identical properties in this assay (not depicted), establishing that behavior in this assay is not dictated by the species of origin. Purified heterotypic complexes were subjected to cross-linking of lysine side chains with Bis(sulfosuccinimidyl)suberate (BS3), followed by electrophoretic separation of products via SDS-PAGE. Cross-linked hK14/mK6 and mK17/mK6 migrated predominantly as tetramers (>40% tetramers, <35% dimers), whereas hK16- and mK16-containing complexes migrated predominantly as dimers (<20%; Fig. 1, b and c). K14, K17, and K16 have comparable numbers of lysine residues (23, 22, and 19, respectively), such that side chain availability likely did not influence cross-linking outcome.

The tetramer-destabilizing Pro188 is located in the first subdomain 1B in hK16 protein. We created chimeric cDNA constructs in which the coding sequences for the 1B subdomains in mK16 and mK17 were swapped, and found that again this region of the rod included the determinants underly-

ing this property (Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200408116/DC1). Altogether these data show that the property of forming urea-unstable heterotetramers is conserved owing to distinct biochemical determinants within subdomain 1B of human and mouse K16 proteins.

Given that the previous experiments were performed on recombinant protein, we sought to validate these findings in vivo. We solubilized cultured mouse primary keratinocytes in either 8 M or 6 M urea and separated the lysate on a blue native acrylamide gel, which resolves protein complexes in their native form according to mass. Western blotting of the resolved keratinocyte lysate revealed that K16-containing complexes were smaller than complexes containing the related type I keratins K14 or K17 (Fig. 1 d). Aside from monomers, all bands contain both types I and II keratins (K6 shown, K5 not shown). In another experiment, primary keratinocyte total protein lysates were separated by anion exchange chromatography as in Fig. 1 a. Here, K16 eluted predominantly in the type I monomer and heterodimer peaks, whereas K14 eluted predominantly in the heterotetramer peak (unpublished data). Together, these results suggest that native keratins present in total protein extracts prepared from cells in primary culture behave similarly to purified recombinant proteins.

Type I keratins exhibit a hydrophobic stripe at the surface of coiled-coil heterodimers

Inspection of the 1B subdomain sequence in mK16 does not reveal obvious amino acid candidates likely to disrupt an α-helix and thus explain its inability to form urea-stable tetramers, unlike the prediction for Pro188 in hK16 (Wawersik et al., 1997). We applied structural modeling with the aim of identifying features that would be unique to subdomain 1B in mK16, relative to other type I keratins. We used the structure of cortexillin I as a template for modeling (PDB 1D7M; Burkhard et al., 2000) as done before (Briki et al., 2002; Hess et al., 2002), given that it consists of a 18-heptad repeat-long, uninterrupted, coiled-coil forming domain. After aligning the target and cortexillin I sequences with ClustalX (Thompson et al., 1997), homology model building was performed using default parameters of Modeller6 (Sali and Blundell, 1993). We created several models of the in-register, parallel 1B domain of vimentin and vari-
We expected that Pro188 in hK16 would kink the \( \alpha \)-helical backbone and create a local disturbance at surface of the dimer (Wawersik et al., 1997). Models of the hK16/K6 dimer exhibited a turn in the backbone at Pro188; however, dimer (Wawersik et al., 1997). Models of the hK16/K6 dimer were compared in these dimers, implying that the peptide backbone and create a local disturbance at surface of the materials, except mK17/mK6 keratin pairs. All models aligned well with the overall geometry of cortexillin I and yielded satisfactory evaluations (see Materials and methods), except mK17/mK6x, which showed a slight out of range value for surface hydrophilicity.

We expected that Pro188 in hK16 would kink the \( \alpha \)-helical backbone and create a local disturbance at surface of the dimer (Wawersik et al., 1997). Models of the hK16/K6 dimer indeed exhibited a turn in the backbone at Pro188; however, the \( \alpha \)-helix surrounding the proline was not severely disrupted (Fig. 2 a). The mK16/K6 backbone proved similar to that of hK16/K6 (Fig. 2 a), with a RMSD of 1.14\( \AA \) over the entire 1B subdomain (14 heptads), compared with a RMSD of 2.07\( \AA \) when overlaid with mK17/K6 (Fig. 2 b). RMSD values decreased to 0.68\( \AA \) when five heptads centered around Pro188 were compared in these dimers, implying that the peptide backbones of the helices are structurally very similar. These modeling efforts nevertheless proved useful in that they exposed an intriguing difference when comparing the side chains of hK16/K6, mK17/K6, and mK16/K6 dimers. In mK17, a hydrophobic stripe of four apolar amino acids, located in b and f positions of consecutive heptad repeats, was exposed to the surface of the dimer, rather than buried within (Fig. 2 c). This hydrophobic stripe spans the region where Pro188 is located in hK16. Moreover, a hydrophilic residue, Gln, replaces one of these hydrophobic residues in mK16 (Fig. 2 d). Otherwise, the hydrophobic stripe is evolutionarily conserved in many type I keratins, including vertebrates and cephalochordates (e.g., Branchio-

**Figure 1.** Mouse and human keratin 16 form unstable tetramers with type II K6. (a) Various purified type I keratins (human K14, mouse K17, human K16, mouse K16) were individually mixed with the mouse type II keratin K6\( \beta \) in a 55.45 \( \text{M ratio, applied to a Mono Q anion-exchange chromatography column, and eluted with a gradient of guanidine-HCl. Fractions were analyzed by SDS-PAGE. Monomeric type II keratins elute first, followed by monomeric type I keratins and heterodimers of type I and II keratins (Wawersik et al., 1997). White lines indicate that intervening lanes have been spliced out. (b) Type I-type II heterotypic complexes (see panel a) were chemically cross-linked with BS3. Cross-linked products (4 \( \mu \)g proteins) were resolved on a 4–16% gradient SDS-PAGE and stained with Coomassie blue. Although individual keratins do not cross-link under these conditions (not depicted; Coulombe and Fuchs, 1999), the type I/type II mixes cross-link as oligomers. The migration standards are indicated at left, and oligomeric complex positions of tetramer (T), dimer (D) type II (II M) and type I (I M) monomers are indicated on the right. Dotted boxes indicate region quantitated in c. The presence of multiple bands for each of the cross-linked heterodimer and heterotetramer products corresponds to distinct intramolecular cross-links. For each concentration tested, antibodies to types I and II keratins react with all of these bands (not depicted), reflecting their heterotypic character. K14, K17, and K16 have comparable numbers of lysine residues (23, 22, and 19, respectively), such that side chain availability likely did not influence cross-linking outcome. There is excellent concordance between the chromatography and chemical cross-linking assays. White lines indicate that intervening lanes have been spliced out. (c) Densitometry was performed on the cross-linked products in b and in replicate experiments. The intensities of monomer, dimer, and tetramer in each lane were summed to 100%, and the percentages of tetramers (black box) and dimers (gray box) were graphed (mean ± SEM). (d) Cultured mouse primary keratinocytes were lysed with 8 M or 6 M urea, mixed with Coomassie G250 dye and separated on a blue native gel (5–13% acrylamide without SDS). Western blotting was performed after transfer to membrane and revealed monomer, dimer, and tetramer complexes of the various native keratins. The approximate migration of various recombinant proteins is indicated at the right; however, these marks should not be considered an exact molecular mass. Although a general calibration curve can be calculated for blue native gels, the apparent mass of a specific complex can vary up to 20% due to different solubilization conditions and post-translational modifications (Schagger, 2001). This explanation may also account for the difference in migration of the dimer band in 8 M versus 6 M urea. Alternatively, there may be a keratin binding protein present in 6 M urea that is dissociated in 8 M urea. Note that two different exposures are shown for K6. The darker exposure (K6\( \beta \)) shows that K6 is present in both monomer and heterodimer bands in 8 M urea. Hydrophobic stripe spans the region where Pro188 is located in hK16. Moreover, a hydrophilic residue, Gln, replaces one of these hydrophobic residues in mK16 (Fig. 2 d). Otherwise, the hydrophobic stripe is evolutionarily conserved in many type I keratins, including vertebrates and cephalochordates (e.g., Branchiostoma; Fig. 2 e). The hydrophobic stripe is not conserved in any other IF sequence type, including type II keratins (Fig. 2 e).

**The hydrophobic stripe in coil 1B of type I keratins mediates heterotetramer stability**. Hydrophobic patches on the surface of proteins are thermodynamically unfavorable in aqueous environments; their presence often is indicative of a region of protein–protein or protein–lipid interaction. In this case, the hydrophobic residues are positioned at a potential interface between two dimers within a keratin tetramer. We postulated that the hydrophobic stripe is important for tetramer stability, and that Pro188 in hK16 would disrupt it by reorienting the side chains of the proximal hydrophobic stripe residues. We designed several mutant proteins that would, according to our postulate, either destabilize...
mK17-containing tetramers or stabilize mK16-containing tetramers. We mutated three of the four hydrophobic residues in mK17 to Gln (mK17 L146Q, V150Q, L157Q; mutant named mK17QQQ) or Ala (mK17 L146A, V150A, L157A; mutant named mK17AAA; Fig. 2 f). Glutamine was chosen because it is the amino acid found in the midst of mK16’s hydrophobic stripe, and it retains the side chain length of the Leu residues it replaces. We chose Ala as a more moderate substitution that maintains the apolar nature of the stripe, but with minimal side chain length. We also swapped four residues between mK16 and mK17 including K16 Q179V and the 3 aa surrounding Pro188 in hK16 (mK16 183QFT to SIL). The reverse was done to K17 (V150Q, 154SIL to QFT). These mutants were designated mK16VSIL and mK17QQFT, respectively (Fig. 2 f).

When assessed by anion exchange chromatography and chemical cross-linking, all three mutant K17/K6 complexes formed unstable tetramers in a similar manner to mK16/K6 (15% tetramers in 8 M urea). In stark contrast, the mK16VSIL/K6 formed stable tetramers similar to mK17/K6 (>35% tetramers in 8 M urea; Fig. 3, a and b). Of note, mK16 Gln179 (the second amino acid in the hydrophobic stripe; Fig. 2, d and e) is conserved in rat K16 (not depicted). Next, we determined that all mutants formed filaments with the same efficiency as wild-type proteins based on a pelleting assay with high speed centrifugation (Fig. 3 c). Analysis of filament morphology using negative staining and electron microscopy revealed that all mutants formed smooth-surfaced, long filaments that appeared similar to their respective wild-type filaments (Fig. 3 e).

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The hydrophobic stripe on type I keratins is accessible on the surface of keratin dimers. We performed binding assays with the hydrophobic-binding fluorophore 8-anilino-1-naphthalenesulfonic acid (ANS), which binds partially folded proteins having small hydrophobic regions exposed to the surface (Semisotnov et al., 1991; Fig. S2, available at http://www.jcb.org/cgi/content/full/jcb.200408116/DC1). Binding of ANS to mK17/K6 heterodimers can reproducibly be detected, and decreases significantly as tetramer formation is allowed to proceed. ANS binding to mK16/K6, mK17AAA/K6, and mK17QQQ/K6 heterodimers is significantly decreased.

The hydrophobic stripe on type I keratins is accessible on the surface of keratin dimers.
significantly lesser than to mK17/K6 or mK16VSIL/K6 heterodimers (Fig. S2). ANS binding studies thus provide direct support for the existence of a hydrophobic stripe contributed by type I keratins at the surface of keratin heterodimers, and reveal a tight correlation between binding and ability to form urea-stable tetramers.

Probing the significance of keratin tetramer stability

We previously showed that K16 could not compete effectively with K14 in the formation of stable heterotypic complexes in the presence of substoichiometric amounts of type II binding partners in vitro (K5 or K6; Paladini et al., 1996). The availability of suitable mouse models created an opportunity to monitor the fate of K16 protein under conditions of limited partner availability (K6 null mice; Wong et al., 2000) or loss of a major type I keratin competitor (K17 null mice; McGowan et al., 2002). Relative to wild-type control, the steady-state levels of K16 protein (but not mRNA) are much decreased in K6 null keratinocytes in primary culture, in which K5 is the only type II keratin left and whose protein level does not become elevated (Fig. 4 a; Wong et al., 2000; Wong and Coulombe, 2003). In contrast, the levels of K14 and K17 proteins are not elevated (Fig. 4 a; Wong et al., 2000; Wong and Coulombe, 2003). Conversely, the levels of K16 protein, but not K14, are altered in these cells (Wong et al., 2000; Wong and Coulombe, 2003). Relative to wild-type control, the steady-state levels of K16 protein (but not mRNA) are much decreased in K6 null keratinocytes in primary culture, in which K5 is the only type II keratin left and whose protein level does not become elevated (Fig. 4 a; Wong et al., 2000; Wong and Coulombe, 2003).

Lack of a pairing partner causes individual keratins to turn over rapidly in cultured cells (Kulesh et al., 1989; Lersch et al., 1989). Given that tetramer subunits comprise the small soluble pool of cytoplasmic IF proteins (Soellner et al., 1985; Chou et al., 1993), we questioned whether the selective loss of K16 protein (Fig. 4 a) reflected an enhanced turnover rate. [35S]Met/Cys pulse-chase experiments performed in K6-null and wild-type keratinocytes showed that K16 protein turned over almost twice as quickly in K6-null compared with wild type (Fig. 4 b). In contrast, K17 protein turnover exhibited no difference between K6-null and wild type (Fig. 4 b). The rate of K14 turnover was slightly enhanced in K6 null compared with wild-type keratinocytes, but less significantly than K16 (Fig. 4 b, 48-h time point). As an additional point, K16 immunoprecipitates showed a lesser amount of type II keratins compared with K14 and K17 immunoprecipitates (Fig. 4 b), suggesting that K16 may be part of smaller heterotypic complexes. K16 was largely excluded from both K14 and K17 immunoprecipitates, whereas sizable amounts of K14 were found in K16 immunoprecipitates (Fig. 4 b).

As previously observed, the affinity of K16 for K5 and K6 is practically identical, and the presence of K14 enhances the likelihood of retrieving K16 in urea-stable heterotetramers in vitro (Paladini et al., 1996). Thus, K16 exhibits a faster turnover rate than the related type I keratins K14 and K17 in a physiological context where type II keratin partners are limiting. Also, K16 is recruited into a distinct subset of keratin heterotypic complexes in living cells. It appears likely that the mixing of K14 and K16, but not K17, within the same soluble subunits could play a role in the intermediate half-life exhibited by K14 in K6 null keratinocytes (Fig. 4 b).
Discussion

Differential keratin tetramer stability and its potential significance in vivo

The ability to obtain high yields of type I/II keratin heterotetramers in the presence of 8 M urea underscores the unusual stability of these complexes, and hints at a key role for hydrophobic interactions during their formation. We identified a novel determinant—a short stripe of four hydrophobic residues aligned on the surface of coil 1B in type I keratins engaged in heterodimers—that underlies this property. This stripe is specific to type I keratins capable of forming urea-stable heterotetramers, such as K14, K17 (Wawersik et al., 1997) and K18 (Yamada et al., 2002), and is imperfectly conserved in K19 (Fradette et al., 1998), K10 (unpublished data), and K16 (Wawersik et al., 1997; this study), all unable to form urea-stable tetramers. Three different assays (anion exchange chromatography, cross-linking, and blue native gel electrophoresis) have shown consistently that K16 forms less stable tetramers than related type I keratins. For mK16, this property is likely due to the discontinuity of the hydrophobic stripe in coil 1B. The duplication of these observations with both purified, recombinant proteins in vitro and native keratin complexes in the context of total protein lysates from cultured primary keratinocytes supports a likely physiological role for this newly defined determinant. The ability to interconvert between urea-stable and urea-unstable heterotetramers through site-directed mutagenesis suggests that this hydrophobic stripe is a key determinant of the differential stability of keratin tetramers under denaturing conditions. Other IFs, including the type III vimentin, are missing this stripe, correlating with the inability of vimentin to form stable tetramers in 6 M urea (Coulombe and Fuchs, 1990). Given that in vitro assembly buffer conditions differ between IF types, it is likely that the driving forces for tetramerization (and further assembly) also differ between IF types. Our report extends others focused on the role of charged residues (Meng et al., 1994; Mehrani et al., 2001), which are periodically distributed in the rod domain (Parry et al., 1977; McLachlan and Stewart, 1982). Charge interactions that strongly influence keratin tetramer stability (without disrupting filament formation) in vitro have been identified in coils 1A, 2A, and 2B, but not in coil 1B (Mehrani et al., 2001). Also, charge interactions between the relatively basic rod-proximal head domain and the relatively acidic rod domain have been shown to influence tetramerization (Hatzfeld and Burba, 1994; Mucke et al., 2004). Comparison of the presence of hydrophobic stripe identified in this work with other hydrophobic and charged residues in the 1B domain reveals an overall negative charge on the COOH-terminal portion of the 1B domain, a more negative charge on the NH2-terminal portion, with the hydrophobic stripe present near the junction of these two domains (Fig. 5, a–d). Here, we provide data that hydrophobic interactions in subdomain 1B are a contributing force in stabilizing keratin tetramers.

Remarkably, distinct amino acids located at discrete positions of coil 1B in human and mouse K16 underlie the formation of less stable tetramers in vitro, raising the prospect of convergent evolution. As further support, the proline residue in human K16 is conserved in both chimp (Acc# XP_511810) and dog K16 (Acc# XP_548101) such that all K16 sequences known to date contain either the proline residue (human, chimp, and dog K16) or an incomplete hydrophobic stripe (rat and mouse K16). A priori, the presence of Pro188 in a “d” position of the heptad repeat in hK16 (normally buried in the coiled-coil hydrophobic core) is at odds with the proposed role for the surface-exposed hydrophobic stripe. We suggest that this Pro residue, which occurs between the third and fourth hy-
Droehun (Fig. 2, a and e), disturbs the local α-helical backbone (as supported by modeling), altering side chain angles in proximal amino acids and in the end, misaligning the surface hydrophobes that normally stabilize keratin dimer–dimer interactions. Such side chain re-orientation can occur in the context of a coiled coil, as previously shown for myosin (Li et al., 2003). The spatial orientation of hydrophobic side-chains within the stripe is likely important in determining their availability for mediating interactions between keratin heterodimers (Er Rafik et al., 2004).

The paucity of knowledge regarding IF assembly in vivo complicates the determination of the significance of differential tetramer stability, observed in an artificial setting in vitro. By the criteria of mild detergent extraction and high speed centrifugation, the soluble pool of IFs in the cytoplasm is small (<1% in keratinocytes) and consists mostly of tetramers (Soellner et al., 1985; Chou et al., 1993). The amount of hK16 retrieved from the soluble pool does not increase when overexpressed in transgenic mouse skin (Paladini and Coulombe, 1999), suggesting that Pro188 does not affect partitioning to the “soluble pool” in living keratinocytes. Tetramer instability could, as shown here, influence protein half-life under specific circumstances defined in part by assembly partner availability. Another possible outcome is an influence on the size or composition of keratin heterotetramers. Our studies in skin keratinocytes in primary culture revealed at least two types of keratin complexes: those rich in K14 and K17; and those rich in K14 and K16 (Fig. 4 b, 0.5-h time point). Monomeric composition can influence the micromechanical properties of keratin filaments in vitro (Yamada et al., 2002). The physiological relevance of these and other possibilities can be addressed in future studies.

Implications for the structure of the polymerization-competent IF tetramer

The discovery of the hydrophobic stripe in the coil 1B domain of type I keratins creates an obvious question: To what does it bind as antiparallel dimers dock alongside one another to form the tetramer? In the answer lies key insight into the axial alignment of antiparallel dimers within the keratin tetramer subunit. Substoichiometric cross-linking of large oligomers of either type I/II or type III IF proteins identified several tetramer conformations by “nearest neighbor analyses” (Geisler et al., 1992; Steinert et al., 1993a,b,c; Mucke et al., 2004). A recent study (Hess et al., 2004) in which site-directed spin labeling and electron paramagnetic resonance (EPR) were used to analyze the interactions occurring as vimentin transitions from a monomeric state to large oligomers in vitro provided direct, site-specific evidence that the A11 intermediate, which places coil 1B of antiparallel dimers en face, corresponds to the earliest tetramer intermediate. The A22 conformation, which places coil 2B of antiparallel dimers en face, occurs concomitant with the formation of larger oligomers (Hess et al., 2004). On one hand, the EPR findings of Hess et al. (2004) yield spatial constraints that are not compatible with the hydrophobic stripe interacting with self in the context of a keratin A11 tetramer. On the other hand, comparative studies based on cross-linking (Steinert et al., 1993c) and mass-per-unit length determination (Herrmann et al., 1999) yielded strong evidence that precise axial stagger of antiparallel dimers differs for keratin and vimentin tetramers.
Higher resolution analyses are needed to identify the site(s) interacting with the newly defined hydrophobic stripe on type I keratins, and the resulting structure of the keratin tetramer subunit. There are ~20 hydrophobic amino acids in coil 1B domain of both types I and II keratins that are not in “a” or “d” heptad positions, and potentially exposed to the surface of the dimer. Short stripes of less than three hydrophobic amino acids exist in the COOH-terminal half of coil 1B in both types I and II keratins (Fig. 5 e) which, depending on their orientation in space, could interact with the hydrophobic stripe identified in the NH2-terminal half of 1B. Whether coiled-coil dimers are supercoiled within tetramers or not (Er Rafik et al., 2004) is an important consideration. As an additional possibility, the hydrophobic stripe could sequentially interact with multiple regions within coil 1B through “axial slippage”, which is believed to occur in several instances. For example, as part of conformational changes within a filament as it reaches its final, most energetically stable structure (Mucke et al., 2004), in response to mechanical stretching (Kreplak et al., 2002) or association with other filaments or interacting proteins (Aebi et al., 1988), or as the basis for the IF-dependent mechanotransduction of signals (Mucke et al., 2004).

Materials and methods

Plasmid generation

The following constructs were used: pET-K14 (Coulombe and Fuchs, 1990), pET-k16, pET-K16b (Paladini et al., 1996), pET-k16b (Bernot et al., 2002). The pET-K17, pET-K16b, pET-K16b plasmids were created by subcloning (New England Biolabs, Inc.) the relevant cDNA (McGowan and Coulombe, 1998) into pET-3d (Studier et al., 1990). The Quick-Change Site-Specific Mutagenesis kit (Stratagene) was used on the pET-K17 and the pET-K16 plasmids with the following sense primers to create the designated mutants (underlined residues are mutant compared with the wild-type sequence): mK17 and the pET-mK16 plasmids with the following sense primers to create the designated mutants (underlined residues are mutant compared with the wild-type sequence): mK17QQQ: 5’TGCAGATTCCTTTGAGATGACGTG3’; mK17QQFT: 5’TGCAGATTCCTTTGAGATGACGTA3’; mK17AAA: 5’TGCAGATTCCTTTGAGAACCG3’; mK179QQ5: 5’TGCAGATTCCTTTGAGATGAC3’; mK169Y9S: 5’TGCAGATTCCTTTGAGATGACA3’.

Recombinant protein purification, anion exchange chromatography, and cross-linking assay

pET plasmids were transformed into BL21(DE3) E. coli for recombinant protein expression. Proteins were extracted from inclusion bodies (Paladini et al., 1996) and solubilized in buffer A containing 6 M urea, 50 mM Tris-HCl, pH 8.0, 1 mM EDTA, 2 mM DTT, and 30 μg/ml PMSF. Proteins were purified to near homogeneity via anion exchange chromatography with a High-Trap Q column (Amersham Biosciences; Wawersik et al., 1997). Purified type I and II keratins (0.5 mg/ml) were mixed (1:5:5 M ratio) for 1 h, loaded onto a Mono Q column (Amersham Biosciences), and eluted with linear gradients of 0–125 mM and 125–300 mM guanidine-HCl in buffer A (Coulombe and Fuchs, 1990; Wawersik et al., 1997). Fractions were analyzed by SDS-PAGE. Heterotypic complexes (200 μg/ml) containing types I and II keratin in a 1:1 M ratio were dialyzed overnight into 25 mM sodium-phosphate buffer, pH 7.4, containing 4, 6, 6.5, 8, or 9 M urea plus 10 mM β-mercaptoethanol. The cross-linker BS2 (Pierce Chemical Co.) was added for 1 h, and products analyzed by SDS-PAGE and gel scanning densitometry.

Blue native gel electrophoresis

A detailed description of blue native gel electrophoresis was provided previously (Schagger, 2001). Primary keratinocytes were cultured and lysed with 8 M or 6 M urea in buffer A (see above). After centrifugation to remove debris, 10 μg lysate was mixed with 5% Coomassie G250 in 500 mM 6-aminohexanoic acid and loaded onto a 5–13% acrylamide gradient gel (48 acrylamide:1.5 bisacrylamide; 25 mM imidazole; 500 mM 6-aminohexanoic acid). After gel electrophoresis, the protein was then transferred onto PVDF membrane (Bio-Rad Laboratories), destained in 30% methanol, 10% acetic acid, and conventional Western analysis was performed.

Filament assembly and pelleting assay

Heterotypic complexes (200 μg/ml) obtained from Mono Q fractionation were denatured in 9 M urea, 1 mM DTT, 25 mM Tris-HCl, pH 7.5, for 4 h. Keratin polymerization was induced by dialysis in 5 mM Tris-HCl, pH 7.5, 1 mM DTT, containing 4 M urea (2 h), 2 M urea (2 h), then no urea for >2 h, all at RT. Polymerized filaments were viewed by negative staining (1% uranyl acetate) on a carbon-coated 400 mesh grid (Ted Pella) with a transmission electron microscope (model CM120; Philips) operated at 60 kV. For the pelleting assay, 50 μl of assembled filaments (~10 μg protein) was subjected to centrifugation in an airfuge at 28 psi for 30 min. The supernatant and pellet were analyzed by SDS-PAGE and gel-scanning densitometry.

Modeling keratin 1B dimers

The crystal structure of Cortexillin I (PCB 1D7M) was chosen as a template for the coiled coil 1B dimer. Alignment of Cortexillin I, vimentin, and keratin sequences was performed using default parameters by CLUSTALX (Thompson et al., 1997). Homology model building was performed using the default parameters for energy minimization of Modeller6 (Sali and Blundell, 1993). No further energy minimization was performed. After a minimum of 10 model-building runs, models of individual keratin heterodimers or vimentin homodimers were almost identical. Models with the lowest energy states were chosen for further analysis. Manual inspection was performed using the Swiss pdb viewer (Guex and Peitsch, 1997), and further evaluation was performed through analyses of 3D profiles (Eisenberg method), atomic interactions (ERRAT), and Ramachandran plots (PROCHECK, all available at UCLA-DOE Institute for Genomics and Proteomics, http://www.doebi.uc.edu/Services/SV/). Variability was assessed by superimposing Cα traces and backbones of select models onto the template and calculating RMSD values for positional differences between equivalent atoms. The protein structures were visualized and analyzed on Molmol (Koradi et al., 1996).

Pulse-chase labeling experiments and immunoprecipitations

Skin keratinocytes were isolated from 3-d-old 129Sv wild-type or K6−/− (Wong et al., 2000) pups and seeded in primary culture in 35- or 60-mm tissue culture plates. After 3 d, or when the plates were 90% confluent, cells were starved with DME medium lacking Met and Cys for 30 min. Cells were then pulse labeled with 0.1 μg/ml [35S]Met/Cys (Easy Tag Express Protein Labeling Mix; PerkinElmer) for 20 min. After washing the cells with PBS, labeled cells were chased with normal medium. After 24, 24, or 48 h, the cells were washed and collected in the presence of protease inhibitors, and stored (~80°C). For immunoprecipitation, cells were solubilized with 2% Empigen BB (Calbiochem) in PBS supplemented with 5 mM EDTA and protease inhibitors. Lysates were analyzed by SDS-PAGE and Western blotting for actin (AC40; Sigma-Aldrich) or β-tubulin (DM1A; Sigma-Aldrich). Protein A Sepharose beads (Amersham Biosciences) were washed with PBS and then bound to rabbit polyclonal antibodies directed against K16 (Bernot et al., 2002), K17 (McGowan and Coulombe, 1998), and K14 (AF 64; Covance). Equal protein amounts from lysates (based on actin or β-tubulin load) were added to conjugated beads and incubated overnight at 4°C. Beads were washed with 0.2% Empigen BB. Bound protein was eluted with 5× gel sample buffer containing β-mercaptoethanol. A second immunoprecipitation was performed on the same lysate to ensure complete epitope(s) deletion. Eluted proteins were subjected to SDS-PAGE and Coomassie staining. The gel was incubated for 20 min in ENLIGHTEN autoradiography enhancer (PerkinElmer), dried, and exposed to BiomaxMR film (Kodak), and analyzed by densitometry.

Online supplemental material

Online supplemental material includes the generation of chimeric proteins in which the 1B subdomain between K17 and K16 has been swapped as well as anion exchange chromatography and cross-linking data obtained using these chimeras (Fig. S1). Also shown is spectroscopy analysis of ANS binding to wild-type and mutant dimers (Fig. S2), which provides direct support for the existence of a hydrophobic stripe contributed by type I keratins at the surface of keratin heterodimers. Online supplement-
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