

## Mechanistic Insights into Phosphoprotein-Binding FHA Domains

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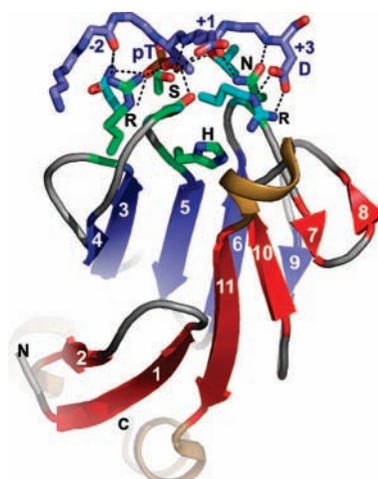
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### CONSPICUOUS

**F**HA domains are protein modules that switch signals in diverse biological pathways by monitoring the phosphorylation of threonine residues of target proteins. As part of the effort to gain insight into cellular avoidance of cancer, FHA domains involved in the cellular response to DNA damage have been especially well-characterized. The complete protein where the FHA domain resides and the interaction partners determine the nature of the signaling. Thus, a key biochemical question is how do FHA domains pick out their partners from among thousands of alternatives in the cell? This Account discusses the structure, affinity, and specificity of FHA domains and the formation of their functional structure.

Although FHA domains share sequence identity at only five loop residues, they all fold into a  $\beta$ -sandwich of two  $\beta$ -sheets. The conserved arginine and serine of the recognition loops recognize the phosphorylation of the threonine targeted. Side chains emanating from loops that join  $\beta$ -strand 4 with 5, 6 with 7, or 10 with 11 make specific contacts with amino acids of the ligand that tailor sequence preferences. Many FHA domains choose a partner in extended conformation, somewhat according to the residue three after the phosphothreonine in sequence (pT + 3 position). One group of FHA domains chooses a short carboxylate-containing side chain at pT + 3. Another group chooses a long, branched aliphatic side chain. A third group prefers other hydrophobic or uncharged polar side chains at pT + 3. However, another FHA domain instead chooses on the basis of pT - 2, pT - 3, and pT + 1 positions. An FHA domain from a marker of human cancer instead chooses a much longer protein fragment that adds a  $\beta$ -strand to its  $\beta$ -sheet and that presents hydrophobic residues from a novel helix to the usual recognition surface. This novel recognition site and more remote sites for the binding of other types of protein partners were predicted for the entire family of FHA domains by a bioinformatics approach.

The phosphopeptide-dependent dynamics of an FHA domain, SH2 domain, and PTB domain suggest a common theme: rigid, preformed binding surfaces support van der Waals contacts that provide favorable binding enthalpy. Despite the lack of pronounced conformational changes in FHA domains linked to binding events, more subtle adjustments may be possible. In the one FHA domain tested, phosphothreonine peptide binding is accompanied by increased flexibility just outside the binding site and increased rigidity across the  $\beta$ -sandwich. The folding of the same FHA domain progresses through near-native intermediates that stabilize the recognition loops in the center of the phosphoprotein-binding surface; this may promote rigidity in the interface and affinity for targets phosphorylated on threonine.



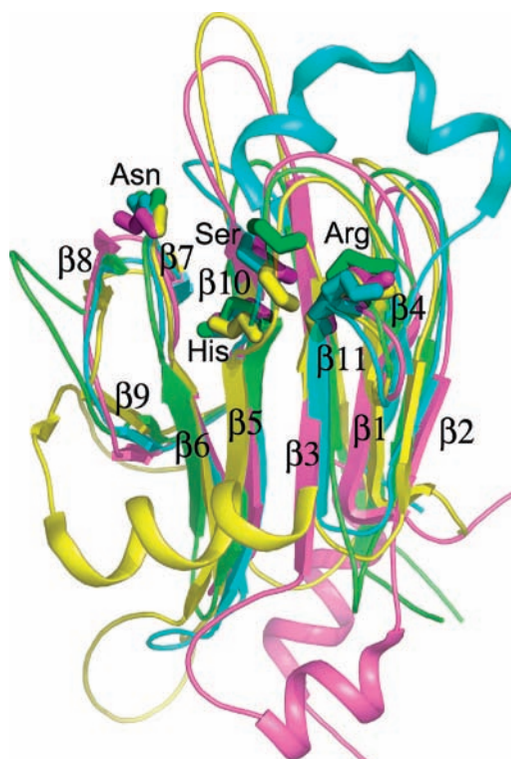
### Introduction

Since phosphate is susceptible to nucleophilic attack, a good leaving group, and retained by cells due to its charge, it is widely exploited as a biological signal.<sup>1,2</sup> Phosphoesters are stable until they meet enzymes such as phosphatases that

rapidly hydrolyze them.<sup>1</sup> Phosphorylation and other post-translational modifications are monitored by noncovalent associations with a diverse array of protein modules that reversibly juxtapose protein partners in intracellular signaling cascades.<sup>3</sup> Since threonine and serine phosphoryla-

tions are the most common protein kinase activities, several classes of protein adapters associate with proteins phosphorylated on hydroxyl-containing threonine residues.<sup>4</sup> These adapters include FHA domains, which assist in regulation of cell growth, division, and differentiation, programmed cell death,<sup>4</sup> DNA damage response, transcription of DNA to RNA, transport of cargo within cells, and protein degradation.<sup>5</sup> FHA domains are found in more than 700 eukaryotic proteins such as kinases, phosphatases, motor proteins called kinesins, DNA-binding transcription factors, RNA-binding proteins, and metabolic enzymes. Identification of the domains in Forkhead transcription factors led to their naming as Forkhead-associated (FHA) domains.<sup>6</sup> FHA domains are found in more than 500 bacterial proteins; some participate in injection of viral proteins into host cells, transmembrane transport, and cell division.<sup>7</sup> A unifying theme among these diverse roles is that FHA domains recognize threonine phosphorylation accompanying activation of protein Ser/Thr kinases.

Since FHA domains are widely important in and representative of biological regulation, this Account focuses on chemical insights into how FHA domains work, starting with their structure, proceeding to specificity, and ending with how the functional structure may form. An FHA domain is organized into a  $\beta$ -sandwich of six-stranded and five-stranded  $\beta$ -sheets (Figure 1). Of the 120–140 amino acids therein, only five are identical in most FHA domains<sup>6</sup> (bold in Figure 2). Four of these are required for phosphoprotein interaction in one example<sup>8</sup> and lie along an edge of the five-stranded  $\beta$ -sheet in three contiguous loops (Figure 1) that contact phosphorylated partners. The central 4–5 recognition loop can be interrupted by a helix<sup>9</sup> (cyan in Figure 1; the loops are numbered by the  $\beta$ -strands they connect). The adjacent 10–11 loop varies in length (top, middle of Figure 1). The FHA domains of Rad53 from baker's yeast have helical insertions in loops distant from the phosphorecognition surface (Figure 1). Structures of FHA domains bound to phosphothreonine (pThr) peptides<sup>9–17</sup> and affinities of pThr peptides for several, varied FHA domains<sup>9,10,12,13,16–18</sup> offer breadth of perspective on the structure and function of diverse FHA domains. General mechanisms shared by FHA domains have emerged: The typically low micromolar dissociation constants imply fast off-rates and depend heavily upon phosphorylation of the target threonine, which enables rapid and reversible switching and dissociation. The five conserved loop residues that support affinity for the phosphorylated threonine (pThr or pT) are surrounded by residues of the loops that differ among FHA domains in order to recognize the preferred sequences. These



**FIGURE 1.** The fold of the FHA domain is illustrated for yeast RAD53 FHA1<sup>11</sup> (magenta) and FHA2<sup>12</sup> (yellow), human Chk2<sup>9</sup> (cyan), and plant kinase-associated protein phosphatase<sup>25</sup> (green). Side chains of five mostly conserved residues near the phosphorecognition surface are drawn.

amino acid positions after or before the pThr are specified by the pT + X or pT – X nomenclature, respectively.

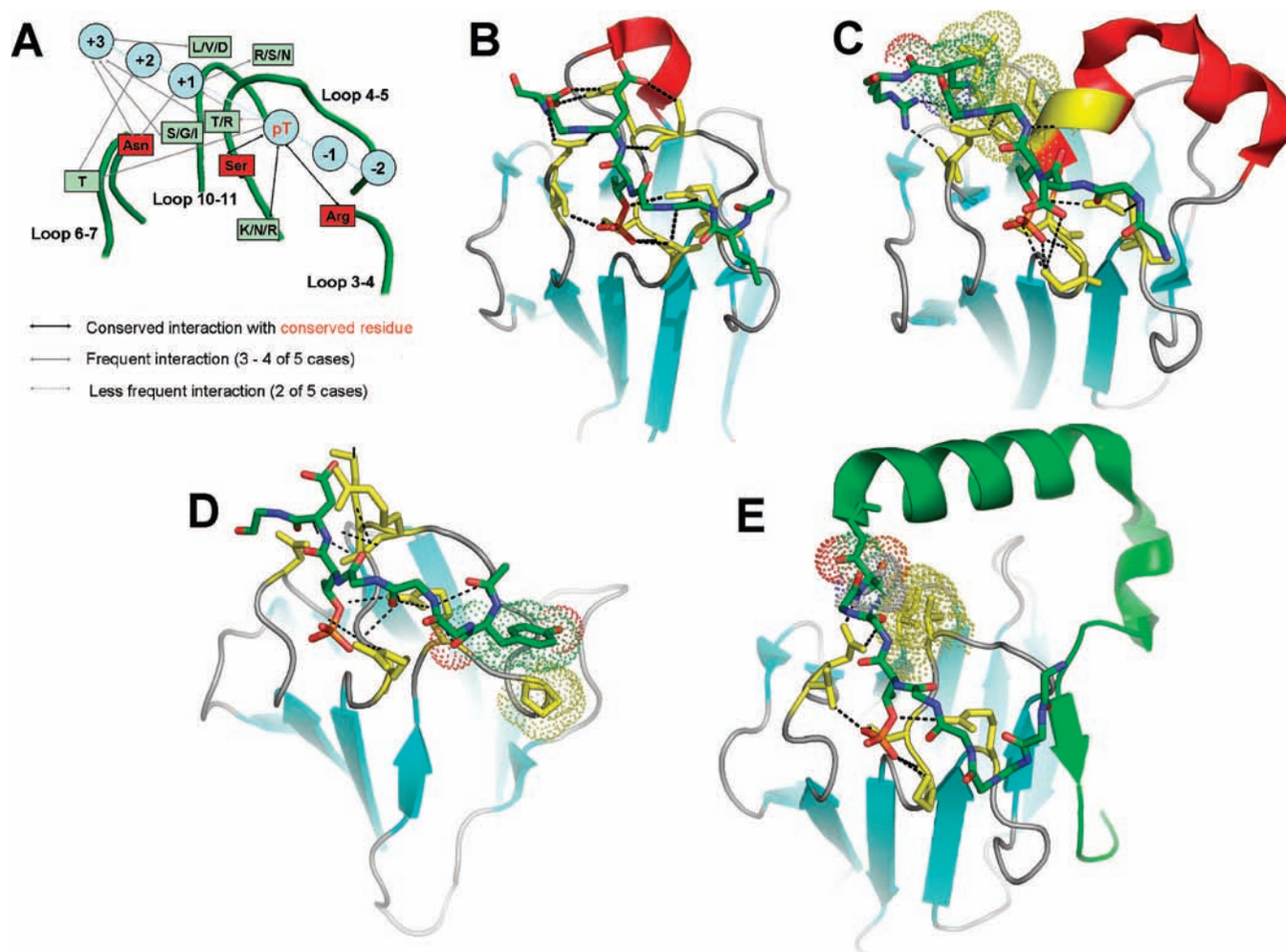
## Determinants of Phosphothreonine Peptide Affinity and Specificity

Residues and interactions shared by FHA domains for recognizing the phosphorylation of the threonine in a target peptide or protein ligand have been identified. Conserved in complexes of phosphopeptides with FHA domains are hydrogen bonds or salt bridges between the phosphate and the conserved arginine of the 3–4 loop, the conserved serine of the 4–5 loop, and a lysine, asparagine, or arginine of the 4–5 loop (Figure 3A). The conserved asparagine of the 6–7 loop often hydrogen bonds the pT + 3 and pT + 1 positions of peptide ligands (Figure 3A–C,E). The conserved glycine and histidine interact with each other and probably stabilize the architecture of the binding site.

Given that phosphorylation of threonine is commonplace in cells and that five-residue protein sequences are enough to encode greater than 100 000 possible phosphothreonine-containing sequences, what sequences does an FHA domain prefer to bind? Distinctive preferences in sequences of phosphothreonine peptides from combinatorial libraries have

	[3-4 lp]	[ 4-5 loop ]	[6-7lp]	[10-11lp]	
Human Ki67	LFGRGI-E-CDIRI----	QL-----P	VVSKQHCKI	8 LHNFSSTNPTQVN	14 VITIIDRSFR-
Human Chfr	TIGRRR-G-CDLSFPSN-----	KL	VSGDHCRI	10 LEDT-STSGTVIN	16 VIYLVYRKN--
human ubiqui ligase RNF8	TVGRGF-G-VTYQLVSKICP----	LM	SRNHCVL	9 IMDNKSLNGVWLN	16 YIQLGVPLENK
Human Chk2	WFGRDK-S-CEYCFDEPLLKRTDKYRTY	<u>S</u> KKHFRI		12 YIEDHSGNGTFVN	16 EIALSLSRN--
Yeast Rad53 FHA1	TFGRNP-A-CDYHLGNI-----	SR	LNKHFQI	8 LLNDISTNGTFLN	16 EITVGVGVES-
Yeast Rad53 FHA2	FIGRSE-D-CNCKI-ED-----	NR	LSRVHCFI	22 WYCHTGTNVSYLN	16 EIKIWDKNNK
Mouse polynucleot kinase	VLGRGP---LTQVTD-----	RK	CSRNOVEL	9 AVKQLGVNPSTVG	16 VLYLVN-----
Arabidopsis KI-FHA	KLGRVS-P-SDLAL----KD----	SE	VSGKHAQI	10 LVDMGSLNGTFLN	24 IITLGTTKVY
Arabidopsis At4g14490	RVGRIVRG-NEIAI----KD----	AG	ISTKHLRI	8 IQDLGSSNGTLLN	16 VIKLGEY----
Human Kinesin C	RVQVD---MDIKLTG-----	QF	IREQHCLF	12 TLEPCEGAETYVN	14 RIVMGKN----
Mouse Afadin 6	EVGTEKFDDNSIQFLG-----	PG	IQPHCDL	8 VTTPRSDAETYVD	14 RLQFGTS----

**FIGURE 2.** Structure-based alignment of sequences of FHA domains. Residues in the 3-4, 4-5, 6-7, and 10-11 recognition loops that contact phosphopeptide ligands are underlined. Mostly conserved residues are bold. Numbers inserted in the sequences count residues omitted from this display emphasizing recognition loops.



**FIGURE 3.** Comparison of structural details of phosphothreonine peptide interactions with FHA domains: (A) summary of FHA domain interactions with peptides for Rad53<sup>11-14</sup> and Chk2,<sup>9</sup> the structures of complexes in each panel represent FHA1 of (B) RAD53 (PDB code 1G6G),<sup>13</sup> (C) Chk2 (1GXG),<sup>9</sup> (D) PNK (1YJM),<sup>17</sup> and (E) KI-67 (2AFF).<sup>16</sup> Each peptide is colored by atom: green for carbon, blue for nitrogen, and red for oxygen. FHA domain residues in contact with ligands are yellow. Electrostatic interactions and hydrogen bonds are shown with dotted lines. Hydrophobic interactions are represented with dots.

been identified for several FHA domains.<sup>13</sup> Both the FHA1 domain of Rad53p from baker's yeast and the FHA domain of Cds1 from fission yeast selectively recognize a pT-V-x-D motif, particularly the negative charge of the aspartate at the pT + 3 position.<sup>13</sup> Three FHA domains tested prefer a hydropho-

bic residue at pT + 3: isoleucine by FHA2 of yeast Rad53p, leucine or isoleucine by human Mdc1, and methionine or tyrosine by Y127 from *Mycobacterium tuberculosis*.<sup>13</sup> The FHA domain of KAPP from plants instead prefers serine or alanine at pT + 3.<sup>13</sup>

### How Does an FHA Domain Achieve Specificity for Its Partners, from among Thousands of Alternative Phosphothreonine Sequences It Might Encounter in the Cell?

As we consider what might adjust the specificity of FHA domains, we next examine shared and varying structural features among complexes of FHA domains with phosphopeptides. The residues FHA domains use to recognize the pT + 3 of ligands vary. Some yeast FHA domains use an arginine to recognize aspartate at pT + 3 in the peptide (for example, Rad53 FHA1), while others select the isoleucine or leucine at pT + 3 with a hydrophobic pocket formed by the 4–5, 6–7, and 10–11 loops. FHA domains usually show weaker selection at the –4 to +2 positions.<sup>13</sup> Contacts between +1 through +3 positions of the phosphopeptide and the 6–7 loop or 10–11 loop of the FHA domain are frequent (Figure 3A). We now examine particular structures of FHA domains with pThr peptides bound, in order to point out interactions that may generalize to other FHA complexes and then proceed in each case to distinctive interactions that may tailor the specificity of that FHA domain.

**Aspartate Preference at pT + 3.** Rad53p regulates the DNA damage response in baker's yeast.<sup>19</sup> In addition to a kinase domain, it comprises FHA1 and FHA2 domains required for DNA damage-dependent Rad53 activation.<sup>20</sup> Its target protein of Rad9 is phosphorylated in response to DNA damage and interacts with the C-terminal FHA domain of Rad53.<sup>21</sup> The interaction of FHA1 of Rad53 and a Rad9p-derived phosphothreonine peptide features an extensive network of hydrogen bonds that align and order the peptide binding loops.<sup>13</sup> A hydrogen bond between the phosphothreonine and the threonine in the 6–7 loop of Rad53 FHA1 (Figure 3B) supplements the conserved contacts with the phosphate (Figure 3A). Side chains of the conserved arginine, conserved asparagine, and Asn86 donate hydrogen bonds to main chain carbonyl oxygen atoms at pT + 1, pT – 2, and pT – 4 positions of the peptide<sup>13</sup> (Figure 3B). The conserved asparagine accepts a H-bond from the amide group of aspartate at pT + 3. These backbone interactions could be widely shared among FHA domain interactions with phosphorylated partners to lend affinity. FHA1 of Rad53 makes other contacts that tailor its distinctive specificity. The carboxylate group of aspartate at pT + 3 in the peptide forms a salt bridge to the guanidinium moiety of Arg83 in the 4–5 loop of FHA1, providing affinity<sup>13</sup> (Figure 3B). Rad53 FHA1 binds to Rad9-derived pThr peptides, containing pTxxD sequences, by contacting pT, pT + 1, and pT + 2 peptide positions using serine and arginine from the 4–5 loop and threonine and asparagine from the 6–7 loop<sup>11</sup> (numbering of secondary struc-

ture in NMR structures of Rad53 domains is one less than here and in other structural reports). Recognition of pThr and aspartate at pT + 3 are keys to the binding affinity and specificity, while pT + 1 and pT + 2 residues appear to fine-tune recognition.

**Preferences for Branched Aliphatics at pT + 3.** FHA2 of Rad53p forms an extensive hydrogen bonding network with a pThr peptide ligand using residues from the 3–4, 4–5, 6–7, and 10–11 loops.<sup>12</sup> The backbone carbonyl oxygen of an asparagine in the 4–5 loop forms a H-bond with the amide proton at pT + 1. A threonine O<sub>γ</sub>H from the 6–7 loop makes a hydrogen bond with the carboxylate group of glutamate at pT + 2.<sup>12</sup> Extensive hydrophobic interactions occur between a threonine methyl group from the 6–7 loop and isoleucine of the 10–11 loop of FHA2 and the peptide's pThr methyl group, glutamate alkyl group at pT + 2, and leucine at pT + 3.<sup>12</sup> The hydrophobic contacts account for FHA2's preference for isoleucine or leucine at the pT + 3 position.<sup>13</sup>

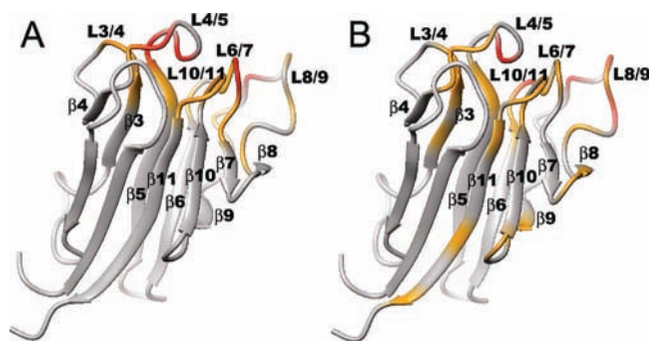
Protein kinase Chk2, mammalian homologue of Rad53p and Cds1 checkpoint kinases from yeasts, is phosphorylated and activated in response to DNA damage, resulting in checkpoint activation and cell cycle arrest.<sup>22</sup> Like FHA2 of Rad53, Chk2 FHA selectively binds pT peptides containing isoleucine or leucine at pT + 3. Hydrophobic residues are preferred at the pT – 2, pT + 1, and pT + 2 positions.<sup>9</sup> The peptide HFDp-TYLI with this sequence pattern packs against the 3–4, 4–5, 6–7, and 10–11 loops of the Chk2 FHA domain.<sup>9</sup> The pThr  $\gamma$ -methyl group makes van der Waals contacts with the conserved asparagine of the 6–7 loop, as well as with threonine, tyrosine, and conserved serine of the 4–5 loop.<sup>9</sup> A lysine in the 4–5 loop forms a salt bridge with the phosphate group and selects for aspartate at pT – 1.

**Changing Preference at pT + 3.** Some FHA domains violate trends in recognition of the key pT + 3 position of the peptide ligand.<sup>20</sup> While FHA1 of Rad53p prefers aspartate at pT + 3,<sup>11,13</sup> it can *switch specificity* to bind a peptide with a physiological recognition site with isoleucine at pT + 3.<sup>14,20</sup> FHA1 interacts similarly with the phosphate of these differing peptides, forming a salt bridge with the conserved arginine of the 3–4 loop and hydrogen bond with the conserved asparagine of the 4–5 loop.<sup>14</sup> FHA1 uses arginine from the 4–5 loop to form a salt bridge with aspartate at pT + 3 of the first peptide. By contrast, FHA1 employs hydrophobic interactions with the pT + 3 position of the physiological peptide sequence; the isoleucine at pT + 3 makes hydrophobic contacts with a glycine and valine from the 10–11 loop of FHA1<sup>14</sup> (Figures 2 and 3A). Moreover, FHA1 makes additional hydrophobic contacts with the physiological site at its pT + 1,

pT + 2, and pT + 4 positions.<sup>14</sup> Thus, FHA1 can recognize pT + 3 with electrostatics or alternatively can recognize pT + 1 through pT + 4 with hydrophobic contacts.<sup>14</sup>

**A Preference for Polar or Hydrophobic Group at pT + 3.** When the kinase-interacting FHA domain (KI-FHA) from plant kinase-associated protein phosphatase recognizes several trans-phosphorylated receptor-like kinase domains,<sup>18,23</sup> it attenuates signaling through receptor-like protein kinases at the cytoplasmic membrane.<sup>24</sup> A principal, but not lone, site of KI-FHA binding *in vitro* was found in BRI1-associated kinase-1 (BAK1) at its Thr546 in region XI near the C-terminal end of the kinase domain.<sup>18</sup> This site was initially suggested by pThr peptides from receptor-like kinase domains that bind KI-FHA. The favorable enthalpy and the hydrophobic effect from contacts with these pThr peptides account for their 8–30  $\mu\text{M}$   $K_D$  values for KI-FHA.<sup>18</sup> The sequence preferences of KI-FHA for pThr peptides<sup>13</sup> are shared with the Thr546 binding site in the BAK1 kinase,<sup>18</sup> namely, the preference for serine at pT + 3 and glutamine at pT + 1. KI-FHA from KAPP is unique in having a long and mobile 8–9 loop that joins the other four loops in recognition of phosphopartners.<sup>18,25,26</sup>

**Choosing Residues N-Terminal to pThr.** An FHA domain with distinctive determinants of specificity has been elucidated.<sup>17</sup> Mammalian polynucleotide kinase (PNK) plays a key role in the repair of DNA with either single-strand breaks or double-strand DNA breaks.<sup>17</sup> PNK acts as a 5'-kinase/3'-phosphatase to create 5'-phosphate/3'-hydroxyl termini required for ligation during repair. PNK is recruited to repair complexes through interactions between its N-terminal FHA domain and phosphorylated components. PNK is notable in not needing contact with the pT + 3 position.<sup>17</sup> Instead, its FHA domain depends more upon pT – 2, pT – 3, and pT + 1 positions of peptide ligands,<sup>27</sup> contrasting other well-studied FHA domains. The equivalent of  $\beta$ -strand 4 in other FHA domains is missing from PNK, but the flanking “3–4” and “4–5” recognition loops remain and form the interface with a peptide ligand.<sup>17</sup> Several electrostatic interactions between a negatively charged pT peptide ligand and the very positively charged surface of the FHA domain of PNK are critical for the micromolar affinity.<sup>17</sup> The second arginine in its “4–5” loop (Figure 2) interacts with the phosphate and aspartate at pT – 3 of the ligand (Figure 3D). The preceding arginine and lysine in the “4–5” loop (Figure 2) interact with the peptide's glutamate at pT – 2 and aspartate at pT + 1. Hydrogen bonds are present between the conserved asparagine and pT + 1 backbone and between the conserved arginine and pT – 2 backbone. Tyrosine at pT – 4 stacks with a proline from PNK (at right in Figure 3D).<sup>17</sup>



**FIGURE 4.** Receptor kinase binding site (panel B) on KI-FHA corresponds to pThr peptide binding site (panel A). Residues with NMR peaks most shifted by phosphorylated partner are colored red and those shifted less yellow. In panel A, the ligand is the pT546 peptide from plant kinase BAK1. In panel B, the partner is the BRI1 kinase domain from plants fused to GST.<sup>18</sup>

## Binding Sites for Phosphoproteins

Phosphothreonine peptides have been essential for studies of associations with FHA domains. Phosphoprotein fragments, however, offer more physiological relevance, but with challenges in obtaining suitable phosphorylation and solubility.

**Similarity of Binding of Globular Phosphoprotein and Phosphopeptide.** BRI1 is a plant receptor Ser/Thr kinase that is critical for brassinosteroid hormonal signaling and development in plants.<sup>28</sup> Once BRI1 is activated, it is attenuated by kinase-associated protein phosphatase (KAPP), which uses its KI-FHA domain to recognize the activated kinase domain.<sup>18</sup> The surface of KI-FHA that binds the 43 kDa kinase domain of BRI1 was mapped by NMR. The BRI1 recognition surface of KI-FHA is essentially same as that for the pThr peptides, involving the 3–4, 4–5, 6–7, 8–9, and 10–11 loops of KI-FHA<sup>18</sup> (Figure 4). The affinity of glutathione-S-transferase (GST)–BRI1 for KI-FHA appears to be moderate, like it is for pThr peptides.<sup>18</sup> Consequently, results on peptide interactions with KI-FHA may offer some value in representing interactions with receptor-like kinases.<sup>18,25,26</sup>

**Examples of Novel Motifs Added to Interface.** Unlike FHA domains discussed above, Ki67FHA lacks affinity for short phosphopeptides. Ki67 is a cancer marker found in the nucleus. The FHA domain at its N-terminus binds human NIFK in a mitosis- and phosphorylation-dependent manner.<sup>29</sup> The structure of the FHA domain of human Ki67 bound to a phosphorylated 44-residue fragment of NIFK is a unique physiological complex.<sup>16</sup> The affinity of the 44-residue hNIFK phosphopeptide for human Ki67FHA is the highest affinity yet demonstrated between an FHA domain and a phosphorylated partner. To gain affinity for Ki67FHA, NIFK must undergo sequential phosphorylation of Thr238 by cyclin-dependent kinase 1 in order for glycogen synthase kinase 3 to phospho-

rylate the Thr234 that Ki67FHA recognizes.<sup>16</sup> When this phosphorylated 44-residue NIFK fragment binds to Ki67FHA, it forms a helix and a  $\beta$ -strand that adds to the edge of the  $\beta$ -sheet of Ki67FHA to form an extended interface that buries 1450 Å<sup>2</sup>.<sup>16</sup> The structure of the recognition surfaces of Ki67FHA is mostly preformed, except that the 1–2, 5–6, and 6–7 loops adjust in conformation when the 44-mer phosphopeptide binds.<sup>16</sup> Three regions of Ki67FHA (Figure 3E) interact with the 44-residue phosphopeptide to provide more contacts and a  $K_D$  of 77 nM,<sup>16</sup> which is about 100-fold greater affinity than typical of short pThr peptides. Three sites contribute to the high affinity between NIFK<sub>226–269</sub>3P and Ki67FHA (Figure 3E). Contributing most to affinity is the phosphorylated central threonine (Thr234) of the NIFK peptide that interacts with the well-characterized pThr binding surface. The phosphate of Thr234 accepts H-bonds donated by the conserved arginine from the 3–4 loop, lysine and conserved serine from the 4–5 loop, and a partly conserved threonine from the 6–7 loop. The backbone of the peptide pT + 1 and pT + 3 positions form hydrogen bonds with the backbone and a side chain, respectively, of the conserved asparagine from the 6–7 loop of Ki67FHA. Next in importance for affinity are NIFK residues 260–264 that form a novel  $\beta$ -strand (green arrow in Figure 3E) hydrogen-bonded antiparallel to  $\beta$ -strand 4.<sup>16</sup> Finally, the 4–5 and 10–11 loops of Ki67FHA dock with the  $\alpha$ -helix formed in the hNIFK peptide once bound. A phenylalanine, leucine, and arginine (side chains not shown) on one side of this helix from NIFK fit into a hydrophobic patch formed by the 4–5 loop and 10–11 loop of Ki67FHA (Figure 3E), likely both stabilizing the helix and enhancing affinity.<sup>16</sup>

Another unique behavior is that the yeast Dun1 FHA domain preferentially binds a sequence having *two* phosphothreonine residues from the Rad53 SQ/TQ cluster domain; this activates the *DUN1*-dependent transcriptional response to DNA damage.<sup>36</sup> The conserved arginine of the 3–4 loop and conserved serine of the 4–5 loop, which in other FHA domains recognize a single phosphothreonine, instead reorient in Dun1 for recognizing the first and second phosphothreonine residues, respectively, from the bound peptide.<sup>36</sup>

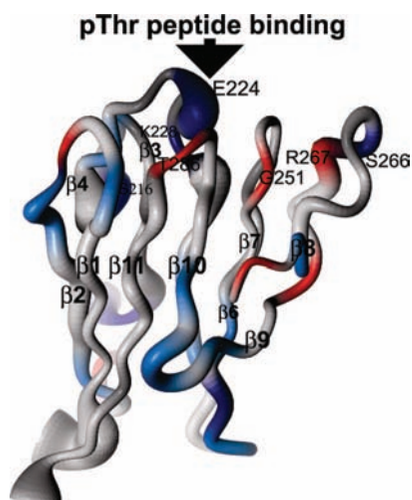
### Other Proposed Protein-Binding Sites

Functional surfaces of FHA domains were predicted<sup>25</sup> by a bioinformatics method.<sup>30</sup> The approach predicts  $\beta$ -strand 4 to be a functional surface, consistent with the structure of Ki67FHA where  $\beta$ -strand 4 is paired with the extended hNIFK peptide<sup>16</sup> (at right in Figure 3E). Since the bioinformatics prediction applies to the entire protein family of FHA domains,<sup>30</sup>  $\beta$ -strand 4 may turn out to contribute to protein recognition events of

other FHA domains as well. The predicted functional surfaces include the phosphoprotein binding surface, broad and variable in properties among FHA domains, and a recognition surface most distant from the phosphoprotein binding site.<sup>25</sup> This patch is centered about Ile157 of Chk2 (at bottom of Figure 3C), includes residues from  $\beta$ -strand 9 and the 5–6 loop,<sup>25</sup> and is the site of the I157T mutation in Li–Fraumeni cancer patients that disrupts binding of BRCA1, CDC25A, and p53 (proteins critically important in cancer).<sup>9</sup> Another functional surface around  $\beta$ -strands 7 and 10 was predicted by the bioinformatics. At this site, the crystalline FHA domain of human CHFr dimerizes such that  $\beta$ 7/8 (a hairpin in all other FHA domain structures) straightens out into single  $\beta$ -strand that H-bonds with the  $\beta$ 7/8 strand from the other chain of the dimer.<sup>31</sup> This swaps strands 9–11 into a  $\beta$ -sandwich with strands 1–7 from the other chain. In KI-FHA,  $\beta$ -strand 8 lacks measurable structural stability<sup>32</sup> and is mobile on the millisecond scale.<sup>26</sup> If low stability and mobility of  $\beta$ -strand 8 generalize to other FHA domains, it could account for  $\beta$ -strand 8 being malleable enough to straighten out in crystals of segment-swapped CHFr.

### Dynamics and Energetics of pThr Peptide Binding

<sup>15</sup>N NMR relaxation of KI-FHA from KAPP indicates that its residues corresponding to those most important for the affinity and specificity of FHA domains are rigid on the fast time scale of picoseconds in KI-FHA.<sup>26</sup> The conserved glycine, arginine, serine, histidine, and asparagine (Figure 1) are rigid in both free and bound states.<sup>26</sup> Of five residues of KI-FHA likely to form the subsite for the pT + 3 residue, a leucine and conserved asparagine of the 6–7 loop and glycine and threonine of the 10–11 loop are rigid.<sup>26</sup> The fifth, a glutamate of the 4–5 loop, becomes rigid only once the pThr peptide binds. The rigidity of these recognition residues confirms that key recognition loops are preformed in KI-FHA. Why might this be? The intrinsically high backbone rigidity of these key residues should diminish free energy costs of loss of conformational entropy upon binding. The rigidity of these key recognition residues is likely to promote enthalpically favorable van der Waals contacts between the pThr peptide and KI-FHA domain. Rigidity promoting such energetically favorable contacts between phosphopeptide and protein was proposed for the recognition of pTyr peptides by an SH2 domain<sup>33</sup> and a PTB domain.<sup>34</sup> Consistent with that outlook, favorable enthalpy of association provides at least half of the free energy of binding of pThr peptides to KI-FHA.<sup>18</sup> Since a rigid, preformed binding site conducive to enthalpically favorable contacts with phosphopeptide is shared among the SH2, PTB,



**FIGURE 5.** Dynamics changes in KI-FHA upon phosphothreonine peptide binding, derived from  $^{15}\text{N}$  NMR relaxation. Blue represents greater rigidity with pThr peptide bound, while orange-red represents mobilization by pThr peptide binding. Dark blue indicates greater rigidification than lighter blue. The diameter of the backbone tube is proportional to greater amplitude of subnanosecond motion in the free state.

and FHA domains tested, this behavior might generalize to other FHA domains as well.

Binding of a pThr peptide increases rigidity of KI-FHA globally. The average rigidity of the backbone on the fast time scale is increased with pThr peptide bound.<sup>26</sup> In fact, sites of binding-enhanced rigidity are observed with the  $\beta$ -sandwich in all four of the  $\beta$ -hairpins (those with blue coloration in Figure 5).<sup>26</sup> The nanosecond fluctuations enriched in its loops are quenched once the phosphopeptide is bound. Perhaps the binding-increased rigidity of KI-FHA reflects the favorable enthalpy of association<sup>18</sup> that might result from improved internal packing and hydrogen bonding.<sup>26</sup>

Some residues of the 3–4, 4–5, and 8–9 recognition loops are comparatively flexible in the free state.<sup>26</sup> The unusual extension, exposure, and nanosecond mobility of the 8–9 loop of KI-FHA might help it adjust to a broader range of receptor-like kinase partners. Flexibility of recognition surfaces of other proteins broadens their specificity while limiting affinity and aiding dissociation.<sup>33,35</sup> The pThr peptide-increased flexibility on the periphery of the binding site<sup>26</sup> could provide part of the favorable entropy of pThr peptide binding to KI-FHA.<sup>18</sup> The solvation entropy gain may overcome entropic costs of the phosphopeptide becoming rigid upon binding.<sup>18</sup>

The dynamics of KI-FHA may also be linked to its stability. At pH 6.3, KI-FHA possesses a network of residues that undergo backbone motions on the millisecond scale (Figure 6).<sup>26</sup> This network traverses  $\beta_2$ ,  $\beta_1$ ,  $\beta_{11}$ , and  $\beta_{10}$  of the six-stranded sheet, plus  $\beta_4$  of the other sheet, where side chains are arrayed in two

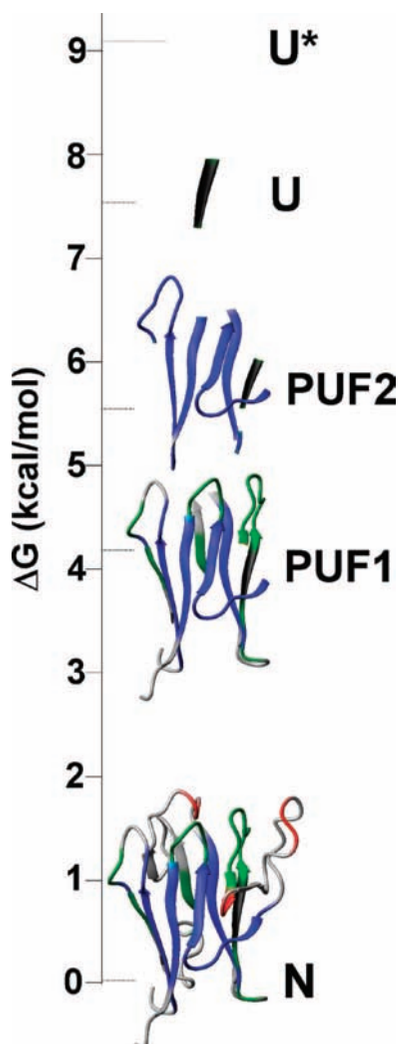


**FIGURE 6.** Sites of slow internal motions of KI-FHA at pH 6.3. Residues with microsecond to millisecond motion at pH 6.3 are colored red where NMR line broadening is larger or orange where it is smaller. This line broadening and motion is quenched at pH 7.3 where KI-FHA is more stable.

packed rows in the interior of the  $\beta$ -sandwich.<sup>26</sup> At pH 7.3 where KI-FHA is at least 3 kcal/mol more stable,<sup>32</sup> the NMR line broadening evidence for the slow motions of this network (Figure 6) vanishes. The stabilization and absence of slow correlated motions at pH 7.3 might reflect improved van der Waals contacts among buried side chains.

### Folding of an FHA Domain and Its Recognition Loops

Compact, near-native intermediates were identified in the folding of KI-FHA.<sup>32</sup> The most stable core of KI-FHA contains  $\beta$ -strands 1, 5, 6, 9, 10, and 11, as well as the 1–2 and 9–10 loops, each minutely populated in the partially unfolded form labeled PUF2.<sup>32</sup> Flanking the most stable core,  $\beta$ -strands 2, 3, and 7, as well as the 6–7 and 10–11 recognition loops, display an intermediate level of stability (green in Figure 7).  $\beta$ -strands 2 and 3 appear to form a subglobal folding motif. Neighboring recognition loops form another subglobal motif of similar stability in PUF1.<sup>32</sup> The least stable subglobal motifs from the 4–5 and 8–9 recognition loops and  $\beta$ -strand 7 appear only in the native state (Figure 7).<sup>32</sup> The near-native intermediate states of KI-FHA suggest that it may fold progressively. An initial hydrophobic collapse may form the stable six-stranded core (PUF2; blue in Figure 7). The 6–7 and 10–11 recognition loops may then organize (PUF1; green in Figure 7). Subsequently, less stable  $\beta$ -strands and loops might add on the flanks of the  $\beta$ -sandwich<sup>32</sup> (Figure 7). Loops of the phosphoprotein binding surface (at top in Figure 7) exhibit strik-



**FIGURE 7.** Hierarchy of folding states of KI-FHA observed under equilibrium conditions. PUF1 and PUF2 are near-native, partially unfolded forms. U is unfolded. U\* has residual structure melted. N is the native state. Stabilities are represented by blue for high global stability, green for intermediate subglobal stability, red for lower subglobal stability, and black for higher-than-global stability.

ing stability absent from the loops on the opposite end of the  $\beta$ -sandwich. The stable formation of the 6–7 and 10–11 recognition loops in PUF1 (green in Figure 7) is significant because (i) these two loops are central to the phosphoprotein binding surface and (ii) they assume considerable stability of 5–6 kcal/mol<sup>32</sup> that may contribute to the aforementioned rigidity of the recognition surface that seems favorable for phosphopeptide binding.

## Conclusions

In representative FHA, SH2, and PTB domains, the subsites important for affinity are rigid. While FHA domains seem to avoid substantial conformational change upon binding, more subtle dynamics changes might accompany their associations

with pThr-containing partners, judging from changes propagating to the periphery of KI-FHA's binding surface and across its  $\beta$ -sandwich. The rigidity expected of preformed phosphothreonine-binding sites of FHA domains could be conducive to favorable enthalpy from van der Waals contacts. The folding of one FHA domain features a near-native intermediate in which the central 6–7 and 10–11 recognition loops are already very stable; this may promote rigidity of the center of the nascent phosphoprotein recognition surface.

A ligand's phosphothreonine is consistently bound by the conserved arginine, serine, and another residue of the 4–5 loop of the FHA domain. Otherwise, FHA domains employ diverse strategies and sequences in recognizing pThr-containing sequences. Dun1 FHA from yeast is distinctive in selectively binding *two* phosphothreonine residues.<sup>36</sup> Three groups of FHA domains select their partners somewhat by the pT + 3 position. Negatively charged aspartate is preferred by yeast Rad53p FHA1 and Cds1.<sup>13</sup> A branched hydrophobic isoleucine or leucine is preferred by Rad53p FHA2 and FHA1, human Chk2, and human Mdc1.<sup>9,13,14</sup> An uncharged polar or hydrophobic residue is preferred by plant KAPP and Y127 from *M. tuberculosis*.<sup>13</sup> Promiscuity by FHA domains is suggested by alternative preferences for aspartate or isoleucine at pT + 3 by FHA1 of Rad53p,<sup>14</sup> acceptance of polar or hydrophobic residue at pT + 3 by KI-FHA and Y127,<sup>13</sup> and PNK's residual preferences.<sup>27</sup> Two important exceptions to pT + 3 recognition have appeared. Mammalian PNK primarily selects for pT – 3, pT – 2, and pT + 1.<sup>27</sup> When these positions are negatively charged, they interact with three positively charged side chains of PNK.<sup>17</sup> Human Ki67FHA instead recognizes in NIFK a phenylalanine from a helix and the addition of a  $\beta$ -strand to  $\beta$ -strand 4<sup>16</sup> (Figure 3E).

Structural contacts that FHA domains use to recognize the key residues listed have been identified but are difficult to categorize since they emanate from variable 4–5, 6–7, and 10–11 loops. Systematic explanation of structural determinants of specificities will require further research. For a greater cross-section of FHA domains, quantitative affinities for phosphorylated partners are needed for describing and differentiating specificities. Since an FHA domain can bind a range of phosphothreonine-containing sequences, additional factors may target it to its intended partners. Explorations of the mechanisms by which more complex, physiological ligands present multiple recognition elements to the FHA domain will be desirable. Additional surfaces of FHA domains may participate in protein interactions. Another frontier is to investigate how FHA domains interact with neighboring modules and how the tandem assemblies jointly recognize their targets in signaling.

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## FOOTNOTES

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