Structure and Location of the Regulatory \( \beta \) Subunits in the \((\alpha\beta\gamma\delta)_4\) Phosphorylase Kinase Complex**

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Background: Structural information concerning the phosphorylatable regulatory \( \beta \) subunit of phosphorylase kinase was lacking.

Results: Chemical, biochemical, biophysical, and computational approaches revealed secondary, tertiary, and quaternary structures for this subunit.

Conclusion: The \( \beta \) subunit is helical and forms the \( \beta_4 \)-bridged core in the \((\alpha\beta\gamma\delta)_4\) kinase complex.

Significance: These findings reveal the architecture of the complex, which provides an explanation for the conformational changes in its bridged core associated with activating \( \beta \)-phosphorylation.

Phosphorylase kinase (PhK) is a hexadecameric \((\alpha\beta\gamma\delta)_4\) complex that regulates glycogenolysis in skeletal muscle. Activity of the catalytic \( \gamma \) subunit is regulated by allosteric activators targeting the regulatory \( \alpha \), \( \beta \), and \( \delta \) subunits. Three-dimensional EM reconstructions of PhK show it to be two large \((\alpha\beta\gamma\delta)_2\) lobes joined with \( D_2 \) symmetry through interconnecting bridges. The subunit composition of these bridges was unknown, although indirect evidence suggested the \( \beta \) subunits may be involved in their formation. We have used biochemical, biophysical, and computational approaches to not only address the quaternary structure of the \( \beta \) subunits within the PhK complex, i.e. whether they compose the bridges, but also their secondary and tertiary structures. The secondary structure of \( \beta \) was determined to be predominantly helical by comparing the CD spectrum of an \( \alpha\beta\gamma\delta_2 \) subcomplex with that of the native \((\alpha\beta\gamma\delta)_4\) complex. An atomic model displaying tertiary structure for the entire \( \beta \) subunit was constructed using chemical cross-linking, MS, threading, and \textit{ab initio} approaches. Nearly all this model is covered by two templates corresponding to glycosyl hydrolase 15 family members and the A subunit of protein phosphatase 2A. Regarding the quaternary structure of the \( \beta \) subunits, they were directly determined to compose the four interconnecting bridges in the \((\alpha\beta\gamma\delta)_4\) kinase core, because a \( \beta_4 \) subcomplex was observed through both chemical cross-linking and top-down MS of PhK. The predicted model of the \( \beta \) subunit was docked within the bridges of a cryo-electron microscopic density envelope of PhK utilizing known surface features of the subunit.

Skeletal muscle phosphorylase kinase (PhK),* a 1.3-MDa \((\alpha\beta\gamma\delta)_4\) complex, catalyzes the \( \text{Ca}^{2+} \)-dependent phosphorylation of glycogen phosphorylase and thereby stimulates the breakdown of glycogen to form glucose 1-phosphate, leading to energy production to help sustain contraction (1). The catalytic \( \gamma \) subunit comprises an N-terminal catalytic core (residues 1–298) that has a typical protein kinase fold (2, 3) and a C-terminal regulatory domain (\( \gamma \)CRD) that binds both the intrinsic calmodulin (\( \delta \)) subunit (4, 5), as well as the regulatory \( \alpha \) subunit (6). The activity of \( \gamma \) in the \((\alpha\beta\gamma\delta)_4\) complex is tightly regulated by the \( \alpha \) (138.4 kDa), \( \beta \) (125.2 kDa), and \( \delta \) (16.7 kDa) subunits, which together promote constraining quaternary (4') interactions on \( \gamma \) (44.7 kDa) that are attenuated by signaling molecules from metabolic, hormonal, and neural pathways that target each of the three regulatory subunits (7, 8). Correspondingly, we have shown that the \( \gamma \)CRD can be chemically cross-linked to \( \alpha \), \( \beta \), and \( \delta \) within the PhK complex (5–7), which led us to postulate that it is the allosteric activation switch through which the three regulatory subunits interact to control the kinase activity of \( \gamma \) (7).

Despite being the first protein kinase to be discovered (9), little is known about the structure of PhK. Reconstructions of images of PhK from EM reveal it to be a \( D_2 \) symmetrical structure of two lobes (10–12), each composed of two \( \alpha\beta\gamma\delta \) protomers packed head-to-head with the lobes associated through interconnecting bridges (13–16). The number of bridges reported for PhK from EM fields of the negatively stained complex on solid grids varies from a minimum of two to a maximum of four (13, 17), with reconstructions of the stained kinase revealing only two large bridges between the lobes (10, 12, 18).

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**β Subunit Structure of Phosphorylase Kinase**

Direct evidence for four bridges was first obtained from a reconstruction of nonactivated PhK from particles of the complex frozen in vitreous ice (cryo-EM), demonstrating that the bridges are viable solution structures and not simply artifacts of staining (11). The cryo-EM results were corroborated by small angle x-ray scattering (SAXS) of nonactivated and Ca\(^{2+}\)-activated PhK (19). SAXS modeling of both conformers demonstrated that only those models containing four, rather than two, bridge structures provided theoretical scattering profiles that accurately fit the experimental data. The divergent results reported above for the bridges in reconstructions of the negatively stained PhK complex may reflect either possible distortions imposed by the mechanical effects of drying and staining on a solid surface or a conformational change associated with the staining process (10). In support of the latter possibility, activation of PhK by Ca\(^{2+}\) has been shown by EM to perturb the structure of the bridges (10) and by SAXS to promote the approach of two bridges toward one another (19). These results suggest that in fully activated forms of PhK, two bridges may abut to form one large bridge, resulting in the overall visualization of only two bridges in low resolution structures of PhK (11, 19).

Attempts to determine the subunit composition of the bridges have proved difficult, because of a lack of structural information for the large homologous α and β subunits, which together constitute over 81% of the total mass of PhK. Although the primary structures are known for both α and β (20, 21), essentially all higher order secondary (2\(^{nd}\)) and tertiary (3\(^{rd}\)) structures reported for these subunits are based on predicted rather than experimental data (22–24). In contrast, high resolution structural information is available for the catalytic core of the γ subunit and for the δ subunit (3, 25); on the other hand, the small size and location of these subunits in the complex make them unlikely candidates for bridging the two (αβγδ)\(_2\) lobes (14, 16). The α subunit is theoretically large enough to span the distances calculated for the bridges (~3 nm); however, its C terminus has been localized to the distal ends of the lobes, which would require the subunit to extend ~90 Å in a straight line through one lobe before traversing the space to the adjacent lobe (10). Moreover, extensive digestion of this subunit in the complex has no effect on the bridges, whereas partial hydrolysis of both lobe (10). Moreover, extensive digestion of this subunit in the complex has no effect on the bridges, whereas partial hydrolysis of both lobes has proved difficult, because of a lack of structural information for the large homologous α and β subunits, which together constitute over 81% of the total mass of PhK. 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Proteolytic mapping of PhK also suggests α to be more peripherally located in the complex than β (26), and localization by immuno-EM of an epitope of the β subunit close to the bridges makes it a more likely candidate for composing the interlobal bridges joining the two (αβγδ)\(_2\) lobes of PhK (16). Nevertheless, no evidence has been previously obtained to indicate the direct involvement of any of the four subunits of PhK in bridge formation.

Functionally, the β subunit places PhK at the interface of metabolic signaling pathways by being the subunit thought to bind the activator ADP (27). It is also the subunit that is predominantly responsible for the regulation of PhK activity by phosphorylation (7). The major regulatory phosphorylatable serine, Ser-26 (28), which is targeted in vivo by cAMP-dependent protein kinase (PKA) and is part of the β subunit’s unique N-terminal phosphorylatable domain (NB1) (20), is also reportedly autophosphorylated, along with Ser-11, by the γ subunit within the complex (7, 20), triggering a conformational change in β that is detected by its cross-linking with 1,5-difluoro-2,4-dinitrobenzene (DFDNB) to form homodimers (29). This possible association of β subunits is consistent with phospho-mimetic S11E/S26E joint mutations that promote self-association of β chimeras in two-hybrid assays (7). We have shown that the NB1 domain is proximal to the γCRD and the C terminus of the β subunit by chemical cross-linking with N-\(\gamma\)-maleimidobutylsuccinimide ester (GMBS) (7). The sum of these data indicates that residues in the β C terminus, the NB1 domain, and γCRD are surface-accessible and proximal to one another within the (αβγδ)\(_4\) complex. Another likely surface-accessible region of β is Ser-700, which is reportedly autophosphorylated by PhK (20). We also report herein an epitope in the PhK complex for an anti-β-specific mAb that has been localized through β-truncation to within a region of this subunit from residues 704 to 815.

The regions of β discussed above represent the little that is experimentally known about its structure and approximate location in low resolution EM structures. A comparison of the sequences of the β subunit and its α subunit homolog indicated that both are multidomain structures, containing subunit-specific regulatory regions and large distinct sequence-similar domains, the latter of which suggested that the subunits are products of early gene duplication events (20). Since these earlier reports, both N-terminal glycosyl hydrolase (GH) clan 15 member-like (GHL) domains and C-terminal calcineurin-B like (CBL) domains have been predicted for α and β (22–24); however, attempts to model the entire structures were unsuccessful (22, 23). We have employed biochemical and biophysical methods in combination with predictive threading and ab initio approaches to model the full β subunit structure as an isolated protein and in the context of the (αβγδ)\(_4\) PhK complex. We report nearly full coverage of the subunit by threading with templates corresponding to GH-15 family members and the protein phosphatase 2A (PP2A) subunit PR65/A (PP2AA). We directly demonstrate for the intact complex by top-down MS and chemical cross-linking that the β subunits compose the central bridge region of PhK and that rigid-body docking of their theoretical counterparts in the individual bridges of the native PhK cryo-EM envelope correlates well with the known structural details for this subunit in the complex (7, 16, 30).

**EXPERIMENTAL PROCEDURES**

**Proteins**—PhK was purified from the psoas muscle of New Zealand White rabbits (31), dialyzed against 50 mM Hepes (pH 6.8), 0.2 mM EDTA, and 10% sucrose, and stored at −80 °C. Its concentration was determined by previously described methods (32). Autophosphorylated PhK was prepared as described previously (33). The mAbs against the α, β, and γ subunits of PhK were previously described (15, 16), and the anti-calmodulin mAb was from Zymed Laboratories Inc.. All other secondary conjugates were from Southern Biotechnology. Truncation mutants of the PhK β subunit were constructed as described previously (7).

**CD**—Far-UV CD spectra were collected for PhK and the αγδ subcomplex using previously described conditions (34). Secondary structure content was estimated using the Dichroweb software package (35), which permits analysis of secondary structure by CONTIN, SELCON, and CDSSTR (36, 37).
Cross-linking—PhK was cross-linked with DFDNB essentially as described (29), with cross-linking initiated by addition of DFDNB and carried out at 30 °C for 2.5 min at pH 8.2 in 50 mM Hepes, 0.2 mM EDTA. Final concentrations of PhK (αβγδε67 monomer) and GMBS in the reaction were 0.47 and 117 μM, respectively. The reaction was terminated by adding an equal volume of SDS buffer (0.125 M Tris (pH 6.8), 20% glycerol, 5% β-mercaptoethanol, 4% SDS), followed by brief vortexing. The PhK sub-units were separated on 6–18% linear gradient polyacrylamide gels and stained with Coomassie Blue. Western blotting of the proteins was performed on PVDF membranes with subunit-specific mAbs as described previously (38). All cross-linking reactions were performed at least twice using different preparations of PhK.

To determine regions of cross-linking in the β monomer, the cross-linked PhK complex was resolved by preparative SDS-PAGE and stained with Coomassie Blue. The bands corresponding to the cross-linked and noncross-linked monomeric β subunits were excised from the gel, sectioned, and exchanged with three aliquots (each ~5× the volume of the gel slice) of 50 mM ammonium bicarbonate, 50% acetonitrile to remove SDS. The proteins were then reduced in 10 mM dithiothreitol for 1 h at 55 °C, and carboxymethylated with 50 mM iodoacetic acid for 1 h in the dark. The gel pieces were washed as described above with 50 mM ammonium bicarbonate, followed by several exchanges with 50 mM ammonium bicarbonate, 50% acetonitrile. After removing the last wash, the gels were dried in a SpeedVac (Savant) and treated with trypsin (Promega; 12.5 ng/μl) for 24 h at 30 °C. Peptides were extracted from the gel pieces with 50% acetonitrile, 5% formic acid.

Bottom-up MS—All samples were concentrated on a CentriVap concentrator (Labconco) to a final volume of 20 μl and pressure-loaded onto a C18 reversed phase nano-column (75-μm inner diameter fused silica packed in-house with 9 cm of 100 Å, 5 μm, Magic C18 particles, Michrom Biosources). Following a wash with 0.1% formic acid for 15 min at 0.5 μl/min, the column was mounted on the electrospray stage of a Fourier transform ion cyclotron resonance MS (LTQ FT, ThermoFinnigan, San Jose, CA), and peptides were eluted at an approximate flow rate of 0.3 μl/min over a 120-min period using a gradient of 0–90% acetonitrile (Buffer A = 0.1% formic acid; Buffer B = acetonitrile, 0.1% formic acid). The source was operated at 1.9 kV, with the ion transfer temperature set to 350 °C. LC MS data were obtained in a hybrid linear ion trap Fourier transform ion cyclotron resonance MS equipped with a 7 tesla magnet. The MS was controlled using an Xcalibur software package to continuously perform mass scan analysis on the FT, followed by MS/MS scans on the ion trap for the six most intense ions, with a dynamic exclusion of two repeat scans (30 s repeat duration and 90 s exclusion duration) of the same ion. Normalized collision energy for MS/MS was set to 35%.

MS Data Analyses—For data analyses, data files were created on Bioworks Browser version 3.2. The corresponding log file was used to generate a list of parent ions for which the corresponding charges and tandem mass spectra were obtained. A list of potential conjugates was generated from the resulting mass list ((M + H)+) by previously described methods (30), and the fragmentation patterns of all viable candidates were analyzed for consistency with the predicted chemistry of cross-linking (30).

The tandem mass spectrum of each conjugate was analyzed using a combination of programs that were subsequently verified by an “in-house” spreadsheet. A list of masses from the tandem mass spectrum of each candidate parent ion was uploaded to MS2Links (39). Theoretical fragmentation of the conjugate in the positive mode was accomplished using the sequences for each peptide in the conjugate and the masses for each intervening chemical cross-link and its possible fragmentation products, using a mass error tolerance not exceeding 0.7 Da for each fragment ion. A modification table was generated for the DFDNB cross-linker based on the loss of NO2, observed for structural analogs of the reagent (40, 41). Potential matching ion assignments for each fragment mass generated by MS2Links were verified using a spreadsheet containing an array of masses predicted for the cross-linker (and fragments thereof) and also those generated for each peptide in the conjugate using MS Product, a fragmentation tool in Protein Prospector developed at the University of California, San Francisco (42). Final assignments were also checked by hand for verification and accuracy.

Top-down MS—A sample (130 μl) of autoprophylated (1.85 and 0.90 mol of P1 incorporated per mol of α and β, respectively) PhK (~1.5 mg/ml) was exchanged into 50 mM ammonium acetate (pH 6.8) using a 5K MWCO VivaSpin column (Vivascience) and then concentrated to a final volume of 30 μl. Non-denaturing nano-electrospray mass spectra were acquired on a Q-TOF 2 mass spectrometer (Micromass/Waters) that was modified for high mass detection (43). Optimal transmission of noncovalent protein complexes was carried out using a previously described protocol (44). Spectra were recorded in positive ion mode with the following settings: capillary voltage 1.7 kV; sample cone 194 V; extraction cone 5 V; collision energy 200 V; collision cell pressure 20 μbar; hexapole ion guide pressure 4.2 × 10−4 μbar; analyzer pressure 1.5 × 10−4 μbar; backing pressure 1.0 × 10⁰ μbar, and TOF pressure 2.1 × 10−6 μbar.

Spectra were acquired and processed using MassLynx version 4.1 (Waters, Manchester, UK) and Analyst QS (MD Science, Applied Biosystems). Errors are reported as ± 1 S.D. All spectra were calibrated externally using a standard (100 mg/ml) solution of cesium iodide, with minimal smoothing and without background subtraction. Subcomplex compositions were determined using the iterative search algorithm SUMMIT (45).

Threading and ab Initio Calculations—The theoretical atomic model of the β subunit was constructed using I-TASSER (46, 47). Using the rabbit muscle β subunit sequence (accession number = P12798) as query, multiple sequence-template alignments were initially generated by the meta-server LOMETS (48), with the domain boundaries decided based on unaligned regions of the threading alignments and the template structures. The β subunit sequence was divided into three domains. The first domain, corresponding to the GHL domain, matched well with several high scoring templates. The full-length model was constructed by I-TASSER, with spatial restraints (Ca distance map and side-chain contacts) extracted from the templates being used to guide replica-exchange Monte Carlo assembly simulation. The second domain, corresponding to the Huntingtin-elongation-A subunit-TOR (HEAT)-repeat like (HRL) domain, had only low scoring templates. The models were then constructed by I-TASSER ab initio structural assem-
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### TABLE 1

Predicted percent secondary structure content of the PhK αγδ-trimer and the PhK hexadecamer from CD analysis at 25 °C and the contributions from individual subunits

| Secondary structure type | Helix | Sheet | Turn | Unordered |
|--------------------------|-------|-------|------|-----------|
| Nonactivated trimer<sup>a</sup> | 65 ± 2 | 14 ± 2 | 5 ± 1 | 16 ± 1 |
| Nonactivated hexadecamer<sup>a</sup> | 64 ± 5 | 16 ± 5 | 6 ± 4 | 14 ± 5 |
| **Trimer subunit** | | | | |
| PhK α | 67 | 16 | 6 | 11 |
| PhK γ<sup>+</sup> | 40 | 18 | 4 | 38 |
| PhK δ | 59 | 12 | 0 | 29 |
| **Hexadecamer subunit** | | | | |
| PhK α | 67 | 16 | 6 | 11 |
| PhK β | 69 | 16 | 8 | 7 |
| PhK γ<sup>+</sup> | 40 | 18 | 4 | 38 |
| PhK δ | 59 | 12 | 0 | 29 |

<sup>a</sup> The percent secondary structure of the PhK holoenzyme and the αγδ-trimer CD spectra was predicted using the CDSSTR algorithm, found in the DICHROWEB software suite (35).

<sup>b</sup> The percent secondary structure of the PhK γ subunit was calculated from the 2.1 Å crystal structure of the PhK γ peptide substrate complex (PDB code 2PHK).

<sup>c</sup> The percent secondary structure of the PhK δ subunit was calculated from the 1.7 Å crystal structure of calmodulin (PDB code 1CLI).

bly, which were guided with the sparse distance and contact restraints from short template fragments (46). The third domain, NB1, was also generated by the I-TASSER ab initio assembly simulations. Decoys generated during the structural assembly simulations were clustered by SPICKER (49), and the cluster centroid models were further refined by REMO to build the full-atomic model (50). The entire β subunit model was constructed by connecting all three domain models together with the domain orientation repacked based on the I-TASSER energy potential. Chemical cross-linking of residues Tyr-51 and Lys-53 from the GHL domain to Lys-1025 from the HRL domain was used as a constraint in packing the domains.

**Rigid Body Docking**—Docking of β atomic models was carried out using MVP-FIT<sup>3</sup>, a program developed by the Zhang laboratory for flexible fitting of atomic models in EM density xmaps. The top 50 atomic models of PhK were docked in the bridge region of the cryo-EM envelope reconstructed from frozen solvated particles of nonactivated PhK (11), based on their localization to this region by top-down MS and intra-subunit chemical cross-linking of β within the PhK complex. The final model was chosen, based on best correspondence with known structural details of β in reconstructions of the PhK complex, as well as best fit in the cryo-EM envelope. All models were visualized using Chimera (51).

### RESULTS

**Determination of 2° Structure of the β Subunit within the PhK Complex**—Predictions of 2° structure for the free β subunit suggest it to be a helical protein (52, 53), with helical content ranging from 41 to 70%. Within the (αβγδ)<sub>4</sub> PhK complex, the structure of β is undoubtedly influenced by subunit interactions with itself and with the other three subunits, as evidenced by chemical cross-linking (7, 29, 30, 38, 54), EM three-dimensional reconstructions (10), and partial proteolysis (26). As opposed to the α, γ, and δ subunits, which have been isolated successfully either individually (8, 55, 56) or in subcomplexes of PhK (57, 58), the β subunit appears to be stable or isolable only in the context of the intact (αβγδ)<sub>4</sub> complex. Although the β subunit undoubtedly influences the structure of α, γ, and δ in the PhK complex, we have shown that features of an established Ca<sup>2+</sup>-dependent communication network among the α, γ, and δ subunits are the same in both the (αβγδ)<sub>4</sub> and (αγδ)<sub>4</sub> complexes (57), suggesting a similar structural architecture for these subunits in each complex. To determine the 2° structure of β within the PhK complex, CD analyses of the intact (αβγδ)<sub>4</sub> complex and an αγδ subcomplex, expressed and purified from insect cells (57), were carried out at 25 °C (Table 1). The percent 2° structural contents (helical, sheet, etc.) of the δ (endogenous calmodulin) subunit (25) and catalytic core of the γ subunit (3) were calculated from known crystal structures of the isolated proteins and were taken to be identical in both the αγδ and (αβγδ)<sub>4</sub> complexes. Differences observed in the CD spectrum for each complex were assumed to reflect differences in contributions from the large homologous α and β subunits, which contribute 74% (α) and 87% (α and β) of the entire 2° structure content of PhK αγδ and (αβγδ)<sub>4</sub>, respectively. Based on 2° structural contributions predicted for α, γ, and δ in each complex using the CDSSTR algorithm (36, 37), the signal difference corresponding to β indicates that it is a helical (69%) protein, with minor amounts of sheet (16%), turn (8%), and unordered (7%) structure.
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PhK complex and/or intramolecular cross-linking of one or both of the β subunits that hinders unfolding of the protein in SDS, leading to an anomalously fast migration (29). To screen for potential intramolecular conjugates, the β monomeric band remaining after treatment with DFDNB was digested in-gel with trypsin, and the resultant peptides were resolved and measured by Fourier transform ion cyclotron resonance MS. The mass list for the digest was then evaluated using the Cross-Search method (30). After culling experimental masses corresponding to those predicted either for nonmodified peptides or for side products of chemical cross-linking, a mass \((m/z_{\text{Exp}} = 2297.207)\) predicted for a peptide corresponding to cross-linking between regions of the N terminus (residues 48–56, LDYYKIVK) and C terminus of β (residues 1021–1028, VDLKLVK) matched that of a peptide mass \((m/z_{\text{Exp}} = 2297.197; 4.37\) ppm error) measured experimentally. The match was verified by MS/MS analysis (Fig. 2), demonstrating the presence of a pool of conjugates with the same peptide partners, but with Lys-1025 in the C-terminal peptide stretch cross-linked to either Tyr-51 or Lys-53 in the N-terminal region of β. No direct evidence for cross-linking between Lys-1025 and Tyr-52 was observed in the MS/MS spectrum. The observed cross-linking between Lys-1025 and both Tyr-51 and Lys-53 within the N-terminal peptide stretch may indicate that these regions may be conformationally dynamic, which would be consistent with the fact that the conformation of β correlates with its phosphorylation and with the activation state of PhK (16, 26, 29). These results with the shorter, rigid cross-linker DFDNB corroborate our previous findings that the N and C termini of β are proximal and provide a viable physical constraint to be used in the three-dimensional modeling of this regulatory subunit.

Three-dimensional models for the β subunit were generated using the I-Tasser hierarchical structural modeling approach (46). Multiple threading alignments of the β sequence were carried out to identify template structures from the Protein Data Bank (PDB) library (48), followed by structural assembly and refinement steps, with subsequent reconstruction of atomic models (50). As shown for the best fit structure (Fig. 3), two major domains comprising residues 41–670 and 671–1092 were predicted and modeled, respectively, using as templates members of the GH family 15 (blue and gray traces) and PP2A (yellow trace). The remainder of the molecule, the small N-terminal phosphoraltatable domain of β (NB1, residues 1–40), is apparently unique and was therefore modeled ab initio (Fig. 3, blue trace) (60). The models were constrained using the DFDNB cross-linking results by forcing the closest possible approach of the α-carbons of Lys-1025 and Tyr-52. In the best fit model, this constraint automatically oriented without further manipulation the side chains of Lys-1025 and Tyr-51/Lys-53 in positions that would allow their observed cross-linking (Fig. 3).

Our multidomain model derived from threading is consistent with previous structural predictions for β that suggested it to be a multidomain structure with an N-terminal GHL domain (22, 24). The GH-15 thread coverage of β is indicated in a two-color scheme, with blue and gray representing, respectively, the \((\alpha/\alpha)_6\)-barrel fold for the catalytic domain (A) and a mixed 2' structure corresponding to the B and C domains found in bacterial and archaeal glucoamylases.
The protein structurally closest to the best model of this region is *Anthrobacter globiformus* glucodextranase (PDB code 1ULV) (61), with a template modeling score of 0.8462 and root mean square deviation of 1.92 Å, indicating a good topographical match (62). The crystal structure of the glucodextranase template revealed the protein in complex with the pseudo tetrasaccharide acarbose, a potent transition state inhibitor of glucoamylases (63). Correspondingly, we recently showed that PhK binds acarbose, which in turn promotes a conformational change in the β subunit (64). The large C-terminal domain of the β model was a relatively good topographical match (template modeling score 0.6272; root mean square deviation 5.41) with PP2A (PDB code 1B3U), and the predominantly helical HEAT repeat structure corresponded well with the helical content estimated for PhKβ by CD measurements (65), revealing an overall helical topology for PhKβ.

**Determination of the 4° Structure of the β Subunits within the PhK Complex**—Having analyzed the 2° and 3° structures of PhKβ, we set out to determine the arrangement of these subunits in the 4° structure of PhK. Both cryo-EM and SAXS analyses of the native (αβγδ)4 PhK complex show it to be a large...
Given the known D2 symmetry through four interconnecting bridges (11, 19), the 4 subcomplex revealed by nano-ESMS of autophosphorylated PhK. Nondenaturing MS of intact phosphorylated PhK yielded two charge state series centered at 12,500 and 16,000 m/z, the latter of which corresponds to the intact hexadecameric PhK complex (discussed in detail in Ref. 72). The former charge state series (blue circles), with charges ranging from +46 to +50, yielded a measured mass of 507,062 ± 55 Da that was in close agreement by mass with only one combination of subunits, namely a β4 tetramer (mass$_{\text{ Theo}}$ = 500,338 Da).

bilateral structure, in which the lobes associate with each other in D2 symmetry through four interconnecting bridges. Previous evidence from our laboratory indicated that an unknown epitope on the β subunit is centrally located on the lobes near the bridges (16) and suggested that at least two β subunits may self-associate (7, 29). If the β subunits actually composed the bridges and self-association were more extensive, we asked whether direct evidence could be obtained for a β4 complex that would by definition represent the bridged central core of PhK upon which the stable αγδ trimers might be arrayed. To investigate this possibility, we first screened the (αβγδ)$_4$ PhK complex with cross-linkers that preferentially targeted the β subunits and analyzed for cross-linked β4 complexes. Several reagents formed conjugates that corresponded to a β4 conjugate by apparent mass and cross-reactivity against subunit-specific mAbs (results not shown). The best resolved species was formed by cross-linking phosphorylated PhK with DFDNB (Fig. 1); however, despite the fact that an anti-α mAb did not detect any α subunit within the β4 band, it did contain minor amounts of co-migrating α/β heteromeric conjugates, because bottom-up MS indicated very small amounts of α peptides in the β4 band. As an alternative corroborating approach, we employed top-down MS to analyze large partially solvated complexes under near-native conditions (66). With the phosphorylated, but not the nonactivated, PhK complex, we observed one subcomplex by nondenaturing nano-electrospray MS that corresponded by mass only to a β4 tetramer (Fig. 4; mass$_{\text{ Theo}}$ = 500.3 kDa; 1.3% difference), with water from the solvation sphere accounting primarily for the small mass addition observed for the tetramer (67). Given the known D2 symmetrical head-to-head packing of αβγδ protomers in each of the two octameric lobes of PhK (15), the only possible arrangement for the β subunits in the complex that can account for these results and our previous findings is a centrally located core, with the four β subunits occupying a region comprising the bridges and internal lobe-faces of PhK.

**DISCUSSION**

Having a three-dimensional cryo-EM envelope for PhK (11), knowledge of the location of the β subunits in its 4° structure, and a model for the β subunit in it 3° structure, we asked whether the modeled β subunits could be docked within the four central bridges of the cryo-EM envelope. To facilitate docking, we reasoned that all known solvent-accessible structures in the modeled β structure, including phosphorylatable serines 11, 26, and 700 (green side chains) and DFDNB-cross-linked residues (red side chains), should be placed on the periphery of the EM structure (Fig. 5A). We also knew the solvent-accessible binding location for the anti-β mAb that was revealed by immuno-EM of PhK in complex with this mAb (16), but we did not know the actual epitope recognized by the mAb. Consequently, we set out to map the epitope by determining the cross-reactivity of the mAb against expressed engineered GST fusion constructs of the complete β subunit and seven previously used truncation mutants as follows: 1–285, 1–350, 1–520, 1–703, 1–815, 1–916, and 33–1092 (7). After these constructs were expressed, purified, resolved on SDS-polyacrylamide gels, transferred to PDF membranes, and probed with the anti-β mAb, only the full-length β and the last three truncation mutants cross-reacted with the mAb (data not shown). Thus, the epitope most likely occurs somewhere in the stretch between residues 704 and 815, and this information was also used in docking the β structure in the PhK EM envelope (Fig. 5A). Only in the orientation shown in Fig. 5B does the shape of the β structure match well with the bridge structure and regions of the lobes, suggesting a β core structure in the complex that encompasses all four bridges. All four subunits were then docked as rigid bodies in the cryo-EM envelope, forming a central core comprising the four bridges and the interior faces of both lobes (Fig. 5C).
with activation in the intact PhK complex (27, 29). NB1 contains the major regulatory phosphorylation site, Ser-26, targeted by PKA in vivo (28), and there have also been several reports that within the PhK complex Ser-11 and Ser-26 are autophosphorylated by the γ subunit (7, 33), leading to activation (28, 33). NB1 is also a potential interaction site for the

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**FIGURE 5. Docking the β model in the PhK cryo-EM envelope.**

A, magnified view of the β subunit docked in the three-dimensional cryo-EM envelope (green mesh) solved for the native PhK complex (11). The β model is shown positioned in the EM envelope, proximal to the interconnecting bridge region of PhK (11), using the known location of an anti-β subunit-specific mAb epitope (mapped to residues 703–815) as a marker (16). All known solvent-accessible structures (phosphorylatable serines 11, 26, and 700; cross-linked residues Lys-1025, Tyr-51, and Lys-53 and the anti-β epitope) are shown located at the periphery of the EM envelope. B, three predominant orientations of PhK are shown, revealing an optimal fit for the β model in the bridge region of PhK. C, β core in PhK, modeled by docking a single color-coded β structure in each of the four bridges of the EM envelope with overall D$_2$ symmetry.
intrinsic calmodulin (δ) subunit, as a synthetic peptide corresponding to residues 5–28 of NB1 inhibited active forms of the PhK complex and had nanomolar affinity for free calmodulin (68). We also previously demonstrated by cross-linking that Arg-18 of NB1 is proximal (±12 Å) to the γCRD (7), which binds the δ subunit within the PhK complex (4, 5). Based on its high affinity for calmodulin, this region of NB1 (residues 5–28) was suggested to have the characteristic amphipathic α-helical structure of many known calmodulin targets (68), an architecture that is also observed in our current ab initio model of this domain. The proximity of NB1 to γ in the PhK complex indicated by cross-linking and by its phosphorylation mandated that this region be surface-accessible and proximal to the position of γ in the EM structure of the (αβγδ)4 complex. The localization of γ was achieved by mapping two-dimensional EM images of PhK in complex with an anti-γ mAb directed against the C-terminal lobe of γ (16). An optimal fit of the β subunit model in the bridge region occurred when satisfying the above conditions.

Aside from the small NB1 domain, the remaining 93% of the 1092-residue β subunit structure was covered by templates from two distinct protein families. The first of these, the GH-15 family of glycosyl hydrolases, provided several good template matches that extended from residues 41 to 670. These results differed from two previous reports that predicted only the GH-15 (α/α) catalytic core to be present in β (22, 24), extending approximately through its first 480 residues (22). Just C-terminal to the GH-15 catalytic core, Carriere et al. (23) predicted a small loop followed by an additional domain (termed B), which extended from residues 492 to 676; however, they were unable to achieve fold recognition for this domain. Our GH fold recognition extended throughout this region and corresponded to the β GH-15 template archaeal glucodextranase B and C domains, which show homology with proteins containing the immunoglobulin folds and carbohydrate-binding domains of several glycosidases, respectively (61). An independent analysis of the primary structure of β using the Pfam database also predicted a GHL fold for this region of the subunit (69). Our predicted domain structures for residues 40–670 of the β subunit mirrored that of the shorter domain predicted by Carriere et al. (23) and are topologically similar to several domains observed in the template glucodextranase crystal structure (61), comprising an (α/α) catalytic core connected to the B and C domains by a short loop (61).

In addition to threading, we used the constraint from intramolecular cross-linking with DFDNB to influence the structure of the β model by positioning the GHL domain proximal to the β C terminus (Fig. 3). In the absence of this constraint, the top 100 models generated for the β subunit did not complement well the shape of the PhK cryo-EM envelope when positioned in all possible orientations in the central bridge region of the molecule. We assume that cross-linking of the β subunit in the intact complex captures a conformation of this subunit that is induced by its interactions with all subunits, including itself, in the 4° structure of PhK. In further support of the use of this cross-linking constraint, we recently demonstrated that PhK binds the glucoamylase inhibitor acarbose and that this binding perturbs intramolecular cross-linking of the β subunit by GMBS (64), which cross-links the GHL domain to the C terminus of the β subunit (30). The direct binding of acarbose by the β GHL domains would allow a potential mechanism for inducing a conformational change in this subunit through altering potential interactions between the predicted β GHL and the HRL domains, consistent with results from the intramolecular cross-linking of β by DFDNB. Acarbose also stimulates the protein kinase activity of PhK γ (64), and functionally it mimics glycogen, which also binds to PhK, activating the catalytic γ subunit (9). It must be cautioned, however, that because the homologous α and β subunits are both predicted to contain GHL domains, we cannot rule out the possibility that effects of acarbose and glycogen on the PhK β and γ subunits are mediated by the binding of these ligands to the α subunit.

Other potential functions for the β subunit have been suggested from models of its C terminus from residues 671 to 1092. From comparing sequence alignments of the α and β subunits from diverse species, Carriere et al. (23) predicted for this region of β (as well as the homologous α) two structurally related α-helical domains as follows: C (residues 711–915) with no fold recognition, and D (residues 915–1092) with a calcineurin B-like (CBL) fold. The CBL fold, identified by hydrophobic cluster analysis (23), is typical of Ca2+-binding proteins containing two globular EF-hand motifs (two perpendicular α-helices tethered by a loop) that are connected by a flexible linker (70). Based on observed chemical cross-linking between the α subunit CBL domain and the catalytic γ subunit (6, 38), the latter of which also binds the EF-hand endogenous CaM (8) subunit (5), it was proposed that the CBL domain targets known CaM-binding sites on γ and/or those predicted for the α and β subunits (23). Although we did not observe a CBL domain on β by threading, the predicted structural relatedness of domains C and D and their predominantly α-helical structure are consistent with our modeling of this region of the subunit as a single domain using the template PP2AA subunit. Crystal structure analysis of PP2AA revealed a molecule in which 15 tandem HEAT repeats (motifs comprising a pair of anti-parallel α-helices) formed a left-handed superhelical structure (65). HEAT-repeat domains have been shown to mediate protein-protein interactions in several proteins, including the PP2AA subunit (65). PP2AA tightly associates with the catalytic PP2A subunit to form a heterodimeric scaffold for binding diverse regulatory subunits that direct the functional phosphatase to specific subcellular sites (71). If present in the β subunit, the HRL domain may function similarly in the context of the PhK complex, forming a scaffold with the GHL domain that promotes interactions with either the catalytic γ subunit or other regulatory β subunits. Such interactions are consistent with the formation of ββ and γγ dimers by cross-linking PhK with DFDNB and GMBS (29, 30), with the observed structural coupling between β and γ in the intact PhK complex (16), and with two-hybrid analyses of β subunit truncation mutants, which demonstrated that the C-terminal half of the HRL domain is required for self-association of the β subunits (7). In addition to a potential correspondence in function, the predominantly helical HEAT-repeat structure agreed well with the 2° structural content estimated for the β subunit and complemented the shape of the bridge structure and interior lobe face after docking β into the PhK cryo-EM envelope.

β Subunit Structure of Phosphorylase Kinase
β Subunit Structure of Phosphorylase Kinase

The first direct evidence for a β₄ subcomplex in the (αβγδ)₄ PhK complex revealed herein by top-down MS and inter-subunit cross-linking of the β subunits by DFDNB provided the rationale for docking these subunits in the central connecting bridge region of the PhK cryo-EM structure. This arrangement of the β subunits is consistent with the observed head-to-head packing of αβγδ protomers in D₄ symmetry (15), which requires any homotetrameric association of subunits to occur at known points of contact between the lobes, i.e. centrally in the complex (Fig. 5). Additionally, our direct results correspond to several reports indirectly linking the β subunits to a central location of the complex by partial proteolysis and EM immuno-localization (16, 17). Taken together, the results herein indicate the presence of a β core in PhK and suggest a structural role for these subunits as a scaffold upon which the α, γ, and δ subunits are arrayed. It is likely that the β subunits position the lobes, and thus the catalytic γ subunits, differently with respect to one another in nonactivated and activated conformers of PhK. For example, we have demonstrated that the conformation of the β subunits correlates with activation of PhK by several mechanisms (10, 26, 29), that the β and γ subunits are structurally coupled to one another during enzyme activation (16), and that changes in the dihedral angles between the lobes and altered bridge structures are readily observed between nonactivated and Ca²⁺-activated forms of PhK by EM and SAXS analyses (10, 19).

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