General and specific promotion of flagellar assembly by a flagellar nucleoside diphosphate kinase

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ABSTRACT Nucleoside diphosphate kinases (NDKs) play a central role in diverse cellular processes using the canonical NDK activity or alternative mechanisms that remain poorly defined. Our study of dimeric NDK5 in a flagellar motility control complex, the radial spoke (RS), has revealed new modalities. The flagella in *Chlamydomonas ndk5* mutant were paralyzed, albeit only deficient in three RS subunits. RS morphology appeared severely changed in averaged cryo-electron tomograms, suggesting that NDK5 is crucial for the intact spokehead formation as well as RS structural stability. Intriguingly, *ndk5*'s flagella were also short, resembling those of an allelic spoke-less mutant. All *ndk5*'s phenotypes were rescued by expressions of NDK5 or a mutated NDK5 lacking the canonical kinase activity. Importantly, the mutated NDK5 that appeared fully functional in *ndk5* cells elicited a dominant-negative effect in wild-type cells, causing paralyzed short flagella with hypophosphorylated, less abundant, but intact RSs, and accumulated hypophosphorylated NDK5 in the cell body. We propose that NDK5 dimer is an RS structural subunit with an additional mechanism that uses cross-talk between the two NDK monomers to accelerate phosphorylation-related assembly of RSs and entire flagella.

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INTRODUCTION

It is well established that nucleoside diphosphate kinases (NDKs) transfer a phosphate from NTPs to NDPs (Berg and Joklik, 1953) via a typical ping-pong mechanism. The γ-phosphate of an NTP first

forms a high-energy intermediate with a conserved histidine (designated as H121 in this article) located in a narrow catalytic cleft. Following the departure of the resulting NDP, the phosphate is transferred to the recipient NDP containing a different base (reviewed by Lascu and Gonin, 2000). These reactions enable the levels of nucleotide species to be appropriately balanced in diverse cellular compartments. Consistent with the fundamental importance in nucleotide metabolism, NDKs are ubiquitous, expressed in both prokaryotes and eukaryotes, often by multiple NDK genes. Based on sequence similarities, NDKs have been divided into group I (NDK1-4) and group II (NDK5-10) (reviewed by Desvignes *et al.*, 2009). NDKs have many aliases derived from independent research, most notably NM23 or NMEs, because of their elevated transcript levels in non- or low metastatic cancer cells (Steeg *et al.*, 1988).

However, accumulating evidence indicates that NDKs are not merely interconverting NDPs and NTPs. They interact with many molecules (reviewed by Steeg *et al.*, 2011). Direct interactions allow NDKs to stimulate GTP loading of their dynamic GTPase partners that drive constant fission and fusion of diverse membranous

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^{*}Address correspondence to: Pinfen Yang (Pinfen.yang@marquette.edu). Abbreviations used: CIAP, calf intestine alkaline phosphatase; DN, dominant negative; HYG, hygromycin; IFT, intraflagellar transport; mAb, monoclonal antibody; MBP, maltose-binding protein; MTDs, microtubule doublets; NDK, nucleoside diphosphate kinase; NG, NeonGreen; P, Paralyzed; PCD, primary ciliary dyskinesia; PMM, paromomycin; RS, radial spoke; S, Swimmers; TAP, Tris-acetate-phosphate; WT, wild type.

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compartments (Boissan et al., 2014). Furthermore, NDK2 modulates G proteins and cation channels as a histidine protein kinase (Srivastava et al., 2006; Di et al., 2010; Cai et al., 2014). It is thought that the phosphate transiently linked to H121 in NDKs can be preferentially transferred to specific residues in target proteins instead of to NDPs (reviewed by Attwood and Wieland, 2015). Although histidine phosphorylation is a common prokaryotic regulatory mechanism, NDK2-mediated histidine phosphorylation is a bona fide eukaryotic strategy for regulating critical processes such as lymphocyte activation. A number of NDK-mediated reactionssuch as modulations of malignancy of cancer cells (e.g., Steeg et al., 1988; MacDonald et al., 1993, 1996; Chang et al., 1996; Wagner et al., 1997; Carotenuto et al., 2013) and transcriptional regulation of c-myc oncogene (e.g., Postel et al., 1993; Thakur et al., 2009)-do not correlate with the NDK activity or require H121. Similarly, a heterozygous NDK mutation (killer-of-prune or kpn) is lethal to a seemingly healthy Drosophila strain with purple eyes (prune) (Sturtevant, 1956; Rosengard et al., 1989; D'Angelo et al., 2004; Carotenuto et al., 2013; Zhang et al., 2015; reviewed by Takacs-Vellai et al., 2015), although the mutation itself does not affect the NDK activity (Lascu et al., 1992). While metastasis suppression and dominant-negative (DN) lethality naturally have attracted much interest for decades, the molecular underpinnings remain uncertain (Steeg et al., 2011).

Group II NDKs diverged from group I before chordate radiation (Desvignes et al., 2009) and perhaps even before the divergence from the last common eukaryotic ancestor. Most group II NDKs have acquired additional molecular modules and primarily reside in motile cilia and flagella, which likely existed in the eukaryotic ancestor (Mitchell, 2007). Flagellar NDKs are built into the microtubule-based axonemal superstructure in motile cilia and flagella (Ogawa et al., 1996; Padma et al., 2001; Munier et al., 2003; Sadek et al., 2003; Patel-King et al., 2004; Duriez et al., 2007). In humans and other mammals, defects in these NDKs results in Primary cilia dyskinesia (PCD), demonstrating their importance in cilia biology (Duriez et al., 2007; Vogel et al., 2012). However, surprisingly, the H121 residue that is required for phosphate transfer between NDPs and NTPs and conserved in group I NDKs is not strictly conserved in group II NDKs (Munier et al., 2003; Patel-King et al., 2004). Furthermore, respiratory cilia of NDK8 PCD patients lacked the NDK8-containing outer dynein complex (Duriez et al., 2007).

These observations led to a prediction that flagellar NDKs are structural proteins and that the H121 in some NDK domains, and thus the NDK activity, is merely an evolutionary relict (Munier et al., 2003). However, other lines of evidence argue that the NDK activity is important. For example, NDK7 copurifies with the γ -tubulin ring complex from the centrosome that supports ciliogenesis at the interphase, and only recombinant NDK7 with H121 could promote microtubule polymerization from the centrosome in vitro (Liu et al., 2014). NDK5 (alias NME5, NDPK5, NM23-5, and RSP23) is a constitutive subunit of the radial spoke (RS), a molecular complex that exhibits NDK activity (Patel-King et al., 2004) and controls flagellar motility. Mouse NDK5 mutation affects the motility of sperm flagella and of both airway and ependymal cilia but does not disturb leftright asymmetry, a process that requires nodal cilia that lack radial spokes (Vogel et al. 2012). Although ciliary and flagellar dyskinesia is the only known phenotype from RS deficiencies in diverse organisms (e.g., Huang et al., 1981; Jeanson et al., 2015), ndk5 knockout mice are sterile and have sperm that are apoptotic, with short flagella or completely aflagellate (Vogel et al., 2012), similar to mutant mice defective in adenylate kinase 7, which is supposed to produce NTP (Fernandez-Gonzales et al., 2009). These observations raised

the possibility that flagellar NDKs are part of a redundant energy metabolic circuitry built into the microtubule-based scaffold to supply NTP for reactions occurring throughout these lengthy organelles (Mitchell *et al.*, 2005; Takei *et al.*, 2014). To elucidate the role of NDK5 in cilia and flagella, we took advantage of in vivo approaches that are uniquely possible in *Chlamydomonas* (Merchant *et al.*, 2007). The results revealed unexpected actions of NDK5 that shed light on NDKs and flagellar biology.

RESULTS

NDK5_{H121A} minigene caused paralyzed short flagella

Previous studies demonstrate that isolated RS complexes exhibit NDK activity and that spoke-less mutant axonemes have half of the NDK activity of wild-type (WT) axonemes (Patel-King *et al.*, 2004). To directly test the NDK activity of NDK5 and to overcome the propensity of NDK5 to precipitate (Munier *et al.*, 1998), we expressed in bacteria maltose-binding protein (MBP)-His-NDK5₁₋₂₀₁ (Donnelly *et al.*, 2006) that contains the conserved NDK domain and the Dpy30 domain but lacks the extended C-terminal tail unique to al-gal NDK5. After cleavage of the MBP moiety in bacteria, soluble His-NDK5₁₋₂₀₁ in the extracts was purified by Ni²⁺ affinity chromatography (Figure 1A, left). The NDK activity was measured using a luciferase-based bioluminometric assay (Karamohamed *et al.*, 1999). As expected, addition of purified His-NDK5₁₋₂₀₁ to the reaction mixture elicited 5×10^5 relative light units (RLU), compared with -0.004×10^5 RLU for the His-HSP40 control (Figure 1A, right).

To elucidate the role of NDK5, we perturbed NDK5 genetically. Although mutations of flagellar genes are mostly recessive (Luck et al., 1977), based on the NDK5's Dpy30 dimerization domain (Sivadas et al., 2012) and the DN effect observed for Drosophila NDK_{kpn} and heteromeric oligomers (Wilkie, 1994), we predicted that expression of inactive NDK5 might also elicit DN effects. Therefore we first engineered an NDK5-6His minigene with the regulatory elements of the LC8 gene, which encodes an abundant 10-kDa protein present in many molecular complexes, including RSs (Yang et al., 2009). We also built into the plasmid a paromomycin (PMM)resistance cassette for selecting transformants. To abolish the NDK activity, the codon for H121, which acts as the transient recipient of the NTP γ -phosphate, was mutated to an alanine codon. The NDK5 and NDK5_{H121A} minigenes were transformed into WT cells. All the screened 300 PMM-resistant colonies in the NDK5 control group were indistinguishable from WT cells, indicating that the minigene itself was not harmful. In contrast, four of 300 clones in the ND-K5_{H121A} group (P/S1-4) contained Paralyzed cells and Swimmers. In three of the four P/S strains, the flagella were about one-half of the expected 10–12 μ m (Figure 1, B and C, p < 0.01). Western blots of flagella did not reveal obvious anomalies but confirmed the presence of His-tagged NDK5 and NDK5_{H121A} in the flagella of the randomly selected S strain and two representative P/S strains (Figure 1D). Quantitative analysis showed similar, if not higher, His-tagged NDK5/Total NDK5 ratios in the S strain flagella (Figure 1E). Thus NDK5_{H121A}, rather than the abundance of His-tagged NDK5 proteins, likely accounted for the motility and length phenotype. While these results were consistent with those in the sperm of ndk5 mice, flagellar lengths in Chlamydomonas varied among the strains; in contrast, none of existing RS mutants was known to exhibit such a prominent length phenotype, except for the LC8 mutant (fla14), whose short flagella were attributed to pleiotropic deficiencies in multiple flagellar complexes (Pazour et al., 1998; Yang et al., 2009). We further investigated an ndk5 recessive mutant and generated genomic constructs to elucidate the role of NDK5 and to test the DN effect of NDK5_{H121A} independently.



FIGURE 1: NDK5_{H121A} expressed from a minigene caused paralyzed short flagella in WT cells. (A) Recombinant His-NDK5₁₋₂₀₁ exhibited NDK activity. Coomassie blue-stained SDS-PAGE showed Ni-NTA purification of His-tagged recombinant proteins (left). Pre, extract from bacteria expressing both MBP-His-NDK5₁₋₂₀₁ and the TVMV protease that could cleave off MBP; Post, the flow-through; Elu, the eluate. Identical amounts of proteins were analyzed in triplicate for NDK activity using the luciferase-based bioluminescence assay (right). RLU, relative light unit. (B) Representative transgenic strains from WT cells transformed with the minigene expressing full-length His-tagged NDK5 (left) or NDK5_{H121A} (right). NDK5_{H121A} transformants had short paralyzed flagella. The cells were fixed before light microscopy for length measurement. (C) Average flagellar length (Fla. Length) of transgenic strains expressing NDK5 or NDK5_{H121A}. All strains from the NDK5 group, represented by randomly selected S1, S2, and S3 strains, contained only swimmers (S) like the parental strain. Four strains from the NDK5 $_{H121A}$ groups contained mostly paralyzed cells but with a fraction of swimmers (P/S). Flagella of three of the four P/S strains had shorter flagella than those of the control strain. Asterisk indicates statistically significant differences (Student's t test, p < 0.01, n = 50). (D) An anti-His Western blot revealed His-tagged NDK5 polypeptides in the flagellar samples from randomly selected S strains and representative P/S strains (left, top). Different protein loads indicated the signals were not oversaturated. Both His-tagged and untagged-NDK5 proteins were revealed by NDK5 polyclonal antibody (right, top). Total proteins were revealed by Ponceau red (bottom). Dynein, dynein heavy chains. (E) Plot Lanes analysis in the ImageJ software showed the ratio of His-tagged NDK5 vs. total NDK5 in the 20 µl flagellar samples in the Western blots in D.

Radial spoke defects in paralyzed flagella of an *ndk5* insertional mutant

An *ndk5* mutant was isolated from a large-scale indexed insertional mutagenesis project (Li *et al.*, 2016). A 21–base pair flanking sequence from high-throughput screening suggested that a PMM-resistance DNA fragment was inserted in the second intron of the *NDK5* gene (Figure 2A) in this strain. The strain was paralyzed as expected for RS mutants. The insertion was confirmed by two PCR amplifications of predicted fragments from the genomic DNA using primer pairs that are conducive for either *ndk5* or its parental strain control ("Con." in Figure 2B, left and middle). A nearby region amplified using a primer pair annealing to the middle of the neighboring gene, *SPL4*, which encodes a putative RNA splicing factor (Figure 2, A and B, right), showed that this region was intact in both

strains. The trace amount of full-length NDK5 (Figure 3A and see *Discussion*) suggests that there is no large deletion at the 5' end of the *NDK5* gene in the mutant.

To segregate potential additional mutations that might have been present in *ndk5*, we backcrossed *ndk5* with WT cells. Although *ndk5* mated poorly, we managed to restore mating ability by crosses with a high mating strain. The strict cosegregation of PMM resistance and paralyzed progeny confirm that NDK5 is required for flagellar motility but is not required for mating.

RS deficiencies of ndk5 flagella

Western blots probed with NDK5 polyclonal antibody detected two major ~120-kDa NDK5 bands in control axonemes. They were greatly diminished in both ndk5 and pf14 that lack most RSs (Diener et al., 1993) (Figure 3A). Also diminished were RSP1 located in the spokehead and HSP40 (alias RSP16) that coassembles with NDK5 and RSP2 in the neck region in WT cells. In contrast, RSP2 and the other representative proteins were not significantly affected, including the other four spokehead proteins (RSP4, 6, 9, and 10). Although RSP3, a phosphorylated RS scaffold protein, appeared normal in abundance, a fraction consistently migrated faster. Proteins assayed as additional controls—inner dynein arm p28, outer dynein arm IC1, and CPC1 from the central pair-were present in normal amounts.

The 120-kDa NDK5 doublet bands revealed by the polyclonal antibody and the NDK activity resembled the 120-kDa doublet-spoke protein bands revealed by the monoclonal antibody (mAb) 3G3 and in-gel kinase assays (Williams *et al.*, 1989; Yang *et al.*, 2004). The absence of 3G3-reactive bands in *ndk5* axonemes that contained 120-kDa RSP2 (Figure 3B) indicated that NDK5, and not RSP2, is the 3G3 antigen, and that it can phosphorylate proteins, at least in vitro, as expected for NDKs. To determine whether the double bands were due to phosphorylation, we treated control

axonemes with calf intestine alkaline phosphatase (CIAP). NDK5 in CIAP-treated axonemes comigrated with the lower band of the NDK5 doublet in the sham-treated control (Figure 3C, top), indicating that the upper band was phosphorylated. In contrast, CIAP treatment did not change RSP3 migration (Figure 3C, bottom).

Short flagella of ndk5

We noticed that *ndk5* and paralyzed backcross progeny assembled shorter flagella than controls at log phase (day 3) in Tris-acetatephosphate (TAP) liquid media; at stationary phase, most cells lack flagella. In minimal media that is commonly used in phenotyping flagellar lengths of mutants (Kubo *et al.*, 2015), most *ndk5* cells had flagella suitable for quantitative analysis. Regardless of distribution or averages, *ndk5* flagella at steady state were consistently shorter



FIGURE 2: Characterization of an *ndk5* insertional mutant. (A) Schematic illustrating NDK5 protein and gene organization. NDK5 has an NDK domain followed by a Dpy-30 domain and an extended C-terminal tail containing three calmodulin-binding IQ motifs. The *ndk5* mutant has an insertion of a 2.6-kb PMM-resistance cassette in the second intron. Black box, exon; white box, intron; black line, flanking sequence; arrowhead, the H121 codon; arrows, PCR primers. (B) PCR diagnosis of the insertion mutagenesis using templates prepared from *ndk5* and the parental strain (Con.). A 0.8-kb fragment at the 3' junction was amplified with the P1S and P1As primer pair from the mutant but not from Con. (left). A nearly 2-kb fragment at the 5' region, including the second intron, was amplified with the P2S and P2As primer pair from the control but not from *ndk5* (middle). An ~0.2-kb fragment from the neighboring *SPL4* gene, ~5-kb upstream to the *NDK5* gene, was amplified with the P3S and P3As primer pair from both templates (right).



FIGURE 3: The *ndk5* mutant is defective in RSs and flagellar assembly. (A) Western blot analysis of axonemes revealed the absence of RSP1, NDK5, and HSP40 in the paralyzed *ndk5* flagella. A fraction of the spoke scaffold protein RSP3 migrated slightly faster than that in WT cells (Con.). The other spoke proteins in the head (RSP4/6 and RSP9/10) and neck (RSP2 and RSP5) appeared normal. The axonemal proteins p28, IC1, and CPC1—subunits of inner dynein arms, outer dynein arms, and the central pair, respectively—were used as controls. The spoke-less *pf14* was a negative control. (B) The antigen of mAb 3G3 is NDK5, and not RSP2, which is present in *ndk5* axonemes. (C) NDK5 is a phosphoprotein. NDK5 in axonemes receiving the sham CIAP treatment migrated as double bands. In CIAP-treated axonemes, all NDK5 migrate with the lower band (arrowhead). CIAP treatment did not change the migration of the spoke scaffold protein RSP3. (D) Microscopy (left) and histogram (right) revealed short flagella (Fla.) in *ndk5* cells cultured in minimal media, in contrast to the full-length flagella of the parental strain (Con.). The length phenotype was more pronounced in late log phase (compare day 4 and day 7, *n* = 50). (E) Average flagellar length corresponding to D. Asterisks indicate statistically significant

than the controls (Figure 3, D and E). The difference became exacerbated as cultures progressed toward late log phase, when nutrients were becoming depleted (compare day 4 and day 7). For evaluation of flagellar generation rate, flagella were first excised from log-phase cells by pH shock. An aliquot of cells was fixed periodically afterward to assess regenerated flagella. Compared with the control, *ndk5* cells regrew flagella more slowly, and new flagella remained shorter even after 90 min (Figure 3F, p < 0.01).

Three-dimensional reconstruction of NDK5 axoneme with cryo-electron tomography

To examine the effect of missing NDK5 (RSP23) on structural integrity of the entire spoke complex, we performed cryo-electron tomography and subsequent subtomogram averaging of *ndk5* axoneme (Figure 4). Our initial analysis from the total averages and comparison with the WT and *pf24* structures from previously published data (Pigino *et al.*, 2011; Bui *et al.*, 2012) revealed the major structural distortion at the spokeneck and spokehead regions in the radial spokes of *ndk5* axoneme. Compared with the WT (Figure 4, A and B), the densities in these regions of *ndk5* axoneme (Figure 4, C and D) appeared much more blurred, or

differences (p < 0.01, n = 50). (F) *ndk5* cells are deficient in flagellar regeneration. Flagella were excised from log-phase cells by pH shock and allowed to regenerate. Aliquots of cells were fixed periodically and imaged. Compared with the control, *ndk5* cells regenerated flagella more slowly, and the final lengths were shorter (p < 0.01, n = 20).



FIGURE 4: Structure of radial spokes from WT (A, B) (EMD-2131; Bui et al., 2012), ndk5 mutant (C, D), and pf24 mutant (E, F; Pigino et al., 2011) Chlamydomonas. Density maps were obtained by averaging subtomograms involving the 96-nm periodic unit from cryo-electron tomography and shown as transverse (A, C, E) and longitudinal (B, D, F) sections. Red arrows indicate corresponding positions of the RS neck. The density at the position of the spokehead components, RSP1, 4, 6, 9, and 10, are missing in averaged cryo-electron tomograms from the ndk5 strain (C, D) similar to the pf24 headless mutant (E, F). Scale bar: 24 nm.

nearly missing, similar to the *pf24* spoke headless axoneme (Figure 4, E and F). Overall length of the spokes from the average in *ndk5* (~29 nm) mutant flagella is like the headless RSs of the *pf24* mutant (Pigino *et al.*, 2011) but considerably shorter than WT spokes (~43 nm) (Bui *et al.*, 2012).

To assess the subtle differences of the spokes within the *ndk5* axoneme and explore the cause of the loss of the spokehead and spokeneck density, we have performed a three-dimensional (3D) variance analysis from aligned subvolumes (Figure 5, A and B), which were included in the final average. The microtubule doublets (MTDs) region displayed no variance as expected and served as a positive control of the success of the density normalization process and the following 3D variance analysis procedures. Interestingly enough, lower variance in the spokeneck and spokehead regions compared with that of the stalk (Figure 5B) excludes the possibility that the RS is fully assembled with bifurcated neck and head but

with orientational flexibility with respect to the spoke stalk. Instead, the vast majority of the spokes may have failed assembling the neck and head.

The clear signal of the stalk region in the variance map indicates heterogeneity, which was not evident from the final average (Figure 5A). To shed light onto the heterogeneous stalk region, we have looked at the aligned subaverages from a few MTDs (Figure 5, C–E). Indeed, as an example, one of the subaverages showed slightly tilted stalks with respect to MTDs (compare red dashed lines in Figure 5, C and D), and another one showed a complete absence of RSs (Figure 5E). This infers both conformational and compositional heterogeneity in the stalk region.

Full rescue of ndk5 phenotypes by NDK5 and NDK5_{H121A}

To determine whether the disrupted NDK5 gene caused both paralysis and short flagella, we created plasmid pNDK5 that contained the NDK5 genomic DNA and conferred hygromycin (HYG) resistance. To test whether NDK activity is required for NDK5 function, we further mutated this genomic construct to pNDK5_{H121A}. The two constructs were transformed into ndk5 cells. Surprisingly, both plasmids fully rescued ndk5 with efficiencies expected for single plasmid transformations. In one experiment, the rescue rate was 50% for the pNDK5 group (22 swimmer clones out of 44 PMM-resistant transformants) and 33% for the pNDK5_{H121A} group (17 out of 51 screened). The rates varied in replica experiments but were not statistically different between the two groups. The rescued strains from both groups were indistinguishable from the ndk5 parental strain (Con.) in both motility and flagellar length (Figure 6, A and B). Western blot analysis showed that NDK5 polypeptides, HSP40, and RSP1 in the axonemes of rescued strains were restored (Figure 6C). Similar migration of both NDK5 and NDK5_{H121A} expressed from the transgene indicates that the H121A mutation did not affect NDK5 migration or phosphorylation. Therefore H121 is not required for the function or phosphorylation of NDK5 polypeptides. Note that the NDK5 from transgenes (Figure 6C, triangles) migrated faster than the endogenous NDK5 in the control. The distance varied in each electrophoresis, perhaps due to slightly different acrylamide concentration. The variation was also observed in other strains (Yang et al., 2008) and may be due to differences at the evolutionarily divergent C-terminal sequence that interacts with calmodulin. Usefully, this feature allowed us to distinguish exogenous and endogenous NDK5 polypeptides. The NDK5 minigene also rescued ndk5, but none of the rescue clones contained 100% swimmers, suggesting that the minigene was less efficient than the genomic construct.

We then compared NDK activities of various axonemes of equal weights. The spoke-less pf14, F₀ ndk5, F₂ ndk5, and, importantly, ndk5 rescued with NDK5_{H121A} exhibited similar NDK activities that were roughly one-half lower than the positive control and the transgenic ndk5 rescued with NDK5 (Figure 6D). Therefore H121A mutation indeed abolished the NDK activity of NDK5. Thus the deficiencies in RS composition, motility, and flagellar lengths of the ndk5 strain are caused by diminished NDK5; and NDK activity of NDK5 requires H121 as do other NDKs, but the enzymatic activity is dispensable for its function and phosphorylation. The phenotypes of strains generated in this project are summarized in Table 1. We further performed site-directed mutagenesis on pNDK5_{H121A} to replace residues predicted to bind NTP or catalyze phosphotransfer, including K12A, N118A, R108A, and H55A (Tepper et al., 1994; Tiwari et al., 2004). The four constructs with double mutations still fully rescued ndk5, indicating that the mechanism of NDK5 action has diverged substantially from the primordial NDK catalytic mechanism.



FIGURE 5: Heterogeneity of radial spokes in *ndk5* axonemes. (A) Density map from the total 96 nm–based average shown as longitudinal sections. (B) A corresponding 3D variance map from aligned and normalized particles. Scale bar: 20 nm. The regions with high variance have darker pixel densities. An absence of variance and electron density in the spokehead and spokeneck indicates that the majority of spokes miss the bifurcated neck and head parts. High variance in the spoke stalks infers either orientation or compositional heterogeneity in that region. An absence of variance in individual MTDs serves as a positive control of applied analysis procedures. The representative density map subaveraged from subvolumes reveals typical straight, rigid conformation of spoke stalk with respect to the doublet in MTD 1 (C), in contrast to the subaverage from MTD 2 (D). Red dashed lines are drawn following the spoke-stalk density for the visual representation of the spoke tilt relative to the doublet. The subaverage from MTD 3 (E) shows complete absence of RS assembly. The numbering of the doublets is arbitrary and does not correspond to accepted numbering in the WT axoneme. Scale bar: 24 nm.

We also converted the genomic construct to express fluorescent NDK5 with a tag containing NeonGreen (NG), a monomeric fluorescent protein with spectral properties similar to EGFP but 2.7-fold brighter (Shaner *et al.*, 2013). Fluorescent NDK5 expressed in *ndk5* transgenic cells decorated the entire flagella, as did NG-tagged RSP3 expressed in *pf14* transgenic cells (Figure 6E). The similar location and intensity of NDK5 and RSP3 suggest that each RS contains two NDK5 molecules, as was found for RSP3 and some other RSPs (Kohno *et al.*, 2011; Sivadas *et al.*, 2012; Oda *et al.*, 2014), and suggests that NDK5 in the RS promotes full-length flagella.

Rescue of the short flagella of a spoke-less *pf14* allelic mutant

We have noticed that flagellar lengths of *p*f14 allelic mutants that lack most RSs vary considerably with growth conditions and from strain to strain. We hypothesized that the putative short flagella phenotype of *p*f14 might be obscured by growth conditions and by backcrosses that led to preferential selection of paralyzed *p*f14 progeny with long flagella conferred by other genes. To test this, we cultured three *p*f14 strains—cc1032, cc2496 and cc613—in minimal media. The latter two strains were derived from a single *p*f14 isolate, whereas cc1032 was derived independently (Luck *et al.*, 1977; Diener *et al.*, 1993; see annotation in the *Chlamydomonas* Resource Center [www.chlamycollection.org/]). As expected, all three strains were paralyzed, and flagellar lengths of the cc1032 and cc2496 strains were within the normal range, around 10–11 µm throughout the culture. However, as in *ndk5*, flagella of cc613 were short, 7.32 ± 0.95 µm on day 3 and barely 5 µm on day 5 (Figure 7). Transformation

of an RSP3 genomic construct into cc613 cells rescued paralysis. Importantly, flagellar lengths (as shown in a representative rescued strain cc613::RSP3) nearly doubled and were no longer sensitive to culture durations. Therefore, like *ndk5*, spoke-less mutants are deficient in flagellar generation, but this prominent phenotype has been overlooked due to variations in the genetic background.

DN effect of NDK5_{H121A} genomic DNA

Figure 1 shows that NDK5_{H121A}-His expressed from a minigene in Chlamydomonas had a DN effect as Drosophila NDK_{kpn}. To test this independently, we transformed WT cells with the untagged $pNDK5_{H121A}$ genomic construct. All transgenic clones in the pNDK5 control group were swimmers (S), whereas some clones for the $pNDK5_{H121A}$ group contained paralyzed (P) cells. The percentages of clones with motility phenotypes varied among repeated transformations. In one experiment, 12 out of 44 PMMresistant clones (27%) contained entirely P cells, whereas 1 clone (2%) had both P and S cells (Figure 8A). The flagella of P strains were also short, albeit not as short as ndk5 flagella, and the length phenotype was exacerbated as the cells approached stationary phase (Figure 8, B and C, compare day 3 and day 7). Therefore NDK5_{H121A}—although fully functional when expressed in ndk5—has a DN effect in the presence of

NDK5, regardless of whether the mutant protein is expressed from the minigene or genomic DNA, or with or without a His tag.

Unexpectedly, Western blots of randomly selected strains showed that, unlike ndk5 axonemes, DN axonemes from the two P strains (P1 and P2) actually contained all RSPs tested, including RSP1, HSP40, and fast-migrating $\mathsf{NDK5}_{\mathsf{H121A}}$ from the transgene (Figure 8D, arrowhead). However, all RSPs were evidently less abundant than in controls, while NDK5 and RSP3 appeared less phosphorylated. The loading control was IC140 of the inner dynein 11. To compare the abundance of endogenous and exogenous NDK5, we analyzed the Western blot with the Plot Lanes program after several attempts that failed to fully separate the two populations. The fast-migrating NDK5_{H121A} (below the dashed line in Figure 8D) constituted 34% and 40% of total NDK5 in the flagellar samples from the two P strains (Figure 8E). This fraction was substantially lower for the other strains: 17%, 12%, 19%, and 16% for S1, S2, P/S1, and P/S2 strains, respectively, and 6% and 9% for the control strains. The similar ratios in S strain and P/S strain samples suggest that a threshold level of NDK5_{H121A} is needed to cause paralysis, whereas the level of NDK5_{H121A} is higher to cause paralysis in all cells. The $pNDK5_{H121A}$ -NG that should have been useful to distinguish the two populations cannot elicit DN effects, perhaps due to the NG moiety.

Hypophosphorylated NDK5 in the cell body

RSP3 is hypophosphorylated in preassembled RSs in extracts from WT cell bodies (Qin *et al.*, 2004) and in mutant flagella with reduced amounts of RSs (Gupta *et al.*, 2012). To elucidate the underpinning



FIGURE 6: Genomic DNA expressing NDK5 or NDK5_{H121A} rescued both the paralysis and length phenotypes of *ndk5*. (A) Flagellar-length (Fla. length) distribution showed that the flagella of *ndk5* transgenic strains expressing NDK5 or NDK5_{H121A} were longer than that of *ndk5* but similar to that of the *ndk5* parental strain. Cells cultured in minimal media were fixed at two indicated time points, and the lengths were measured from micrographs; n = 50. (B) Average flagellar length of corresponding data from Figure 4A. Asterisk indicates statistically significant differences (p < 0.01). (C) Western blot analysis of axonemes from representative transformants. RSP1, HSP40, and NDK5 polypeptides were restored. Note that NDK5 and NDK5_{H121A} expressed by the BAC-derived transgene (triangle) migrated identically but faster than endogenous NDK5 (dot). (D) H121A mutation abolished NDK activity. The NDK activity of axonemes from *ndk5* transformants expressing NDK5 was restored to the activity level of WT axonemes (Con.), whereas the activity of axonemes from *ndk5* transformants. RDK5_{H121A} transformants, spoke-less mutant *pf14*, and *ndk5* strains (either F₀ or F₂ from backcross) was ~50% lower. The relative NDK activity was the RLU value normalized to the control. Quadruple aliquots from each sample were measured. Asterisk indicates statistically significant differences (Tukey HSD test, p < 0.01). (E) Live-cell imaging of transgenic *ndk5* and *pf14*, respectively, expressing fluorescent NDK5 (top) and RSP3 (middle). *ndk5* (bottom) was a negative control. NG-tagged NDK5 and RSP3 appeared identical in distribution and intensity throughout flagella.

	WT	ndk5	ndk5::NDK5	ndk5::H ₁₂₁ A	WT::H ₁₂₁ A(P/S)	WT::H ₁₂₁ A(P)
Motility	S	Р	S	S	P/S	Р
Flagellar generation	++	+	++	++	++	+
HSP40; RSP1	++	-	++	++	++	+
NDK5	++	-	++	++	++	+
NDK5 Pho.	++	N/A	++	++	++	+
RSP3	++	++	++	++	++	+
RSP3 Pho.	++	++	++	++	++	+

+, Reduction in flagellar lengths, protein abundance, or phosphorylation (Pho.); ++, similar to the WT strain;

-, absence of proteins; N/A, not applicable; P, paralyzed cells; S, swimmers.

TABLE 1: Strains described in this study and their corresponding motility, flagellar generation, and RS assembly phenotypes.



FIGURE 7: Diminished RSs result in short flagella. (A) Flagellar-length (Fla. length) distribution of three *pf14* strains cultured in minimal media. The defect in the *RSP3* gene of cc613 and cc2496 that were derived from one common isolate was identical. Only flagella of cc613 were particularly short, and this was exacerbated at late log phase (day 5). The length phenotype and paralysis were rescued by a *RSP3* transgene. (B) Average flagellar lengths of *pf14* cells. Asterisks indicate statistically significant differences (p < 0.01, n = 50).

of the DN effects, we analyzed NDK5 in cell body extracts from various strains by Western blot analysis (Figure 9). Axonemes (lanes 1 and 2) and cell body extracts (lanes 5 and 6) of *ndk5* and its parental strain were controls. The lanes loaded with both axonemes and cell body extracts (lanes 3, 4, and 9) tested whether sample preparations affected polypeptide migration. Given reduced RSs in the DN flagella, we also included a *pf14* cell body extract (lane 8) as a control for impaired RS assembly. As cell bodies typically harbor enough axonemal proteins for the regeneration of two half-length flagella (Rosenbaum and Child, 1967), we reasoned that RSs that failed to assemble in flagella may accumulate in the cell body (Diener *et al.*, 2011).

Anti-NDK5₁₋₂₀₁ polyclonal antibody revealed a faint putative NDK5 band in the parental control cell body extracts, which largely comigrated with the hypophosphorylated NDK5 band in control axonemes (Figure 9A, compare dots in lanes 1, 3, and 5). Similar bands were more prominent in cell body extracts of the P1 DN strain that contained both endogenous (dot in lane 7) and exogenous NDK5 (triangle) or of *p*114 that migrated slightly faster (x in lane 8). These bands in the cell body of all strains except *ndk5* were confirmed by probing the duplicated blot with affinity-purified polyclonal anti-NDK5₈₋₅₈₆ (Patel-King *et al.*, 2004) (dot, triangle, and x in Figure 9B, top). Protein loads are shown in the Ponceau S–stained blot (bottom). The 3G3 mAb was too weak to reveal NDK5 in cell

body samples. Duplicated control blots probed for other RSPs contained multiple background bands and were not informative.

NDK5 is primarily hypophosphorylated in the cell body extracts, suggesting that it becomes phosphorylated after entering flagella. The P1 cell body extract contained more NDK5 polypeptides than the control but was similar to *pf14* (Figure 9A, compare lanes 5, 7, and 8), indicating that NDK5_{H121A}, which causes DN effects, is not excessively overexpressed, and is not degraded either, unlike the presumptive effect caused by the *Drosophila kpn* mutation (Biggs *et al.*, 1988). The varied abundance of NDK5_{H121A} expressed by minigene or genomic constructs in individual transformants may account for variations in phenotype severity, a scenario similar to *kpn* mutants (Biggs *et al.*, 1988).

DISCUSSION

NDK plays an important role in multiple biological processes. Only some of them involve the nucleoside phosphorylase activity for which the proteins were named. More uncertain are mechanisms in the events where the NDK activity is dispensable. The ability to manipulate the gene in transgenic strains of *Chlamydomonas* provides a unique opportunity to investigate the NDK activity–independent mechanism and the DN effects of NDK.

The structural roles of NDK5 for proper assembly of the RS head–neck region

While both Western blot and cryo-electron tomographic analysis reveal RS deficiencies in the ndk5 flagella, the severities differ drastically. Except for RSP1, HSP40, and NDK5, all the other components in the head-neck region, appear in normal abundance in Western blots (Figure 3). However, entire spokeheads (Figures 4 and 5) are invisible in the comparative subtomogram studies. These contradictions could be explained by nonmutually exclusive possibilities. The assembly level of partial spokeheads in ndk5 may be affected by culture conditions, as in the spokehead mutant pf26 (Huang et al., 1981; Wei et al., 2010). Early harvest of sufficient ndk5 flagella for Western blots may account for the seemingly normal abundance. Furthermore, moderate reduced abundance may not be revealed unequivocally by Western blots. Or, the mal-assembled components, at least RSP3's C-terminal extension in the spokehead (Oda et al., 2014), may not confer substantial density above the background. Regardless, NDK5 is necessary for the assembly of intact spokes, especially the neck and head regions. Our tomographic data analysis clearly explains why the ndk5 mutants are paralyzed, similar to other headless RS mutants and cilia of PCD patients with the defective RSP1 gene (Lin et al., 2014) and unlike the jerky flagella that lack only HSP40 (Yang et al., 2008).

Therefore NDK5 has two structural effects. Diminished RSP1 and HSP40 in ndk5 flagella (Figure 3) support the predictions that flagellar NDKs are structural proteins (Munier et al., 2003) and RSP1 is assembled differently in comparison with the other four head proteins (Luck et al., 1977) and are consistent with the chemical crosslinking of NDK5 and RSP1 (Kohno et al., 2011). Because NDK5 is present in RSs in mutant flagella missing HSP40 or RSP1 (Patel-King et al., 2004; Yang et al., 2005, 2008), it is NDK5 that tethers RSP1 and HSP40 to the RS scaffold, rather than vice versa. In addition, NDK5 is important for structural integrity of the RS head-neck regions as well as structural stability of the RS stalk. Without NDK5, the rest of the proteins in this region do not form stable bifurcated neck or spokehead, as shown in subtomogram average of ndk5 axonemes (Figure 4). NDK5, probably cooperating with HSP40 and RSP1, plays an essential role for stable formation of the bifurcated neck and spokehead.



FIGURE 8: NDK5_{H121A} expression from a genomic transgene in WT cells impaired flagellar motility, length, RS assembly, and RS phosphorylation. (A) The percentages of clones with swimmers (S), paralyzed cells (P), or a mixture (P/S) expressing NDK5 (first bar) or NDK5_{H121A} (second bar) from one experiment. Average flagellar length (B) and flagellar-length (Fla. length) distribution (C) of representative strains cultured in minimal media showed that flagellar lengths of P strains (P1) were between the control and *ndk5*, whereas P/S strains (P/S1) had flagella of normal length. Asterisks indicate statistically significant differences (p < 0.01, n = 50). (D) Western blot analysis of axonemes from transformants with different motility levels. RSPs in the flagella of P/S strains (P/S1 and P/S2) appeared normal but reduced in P strains (P1 and P2) (top). In addition, NDK5 and the scaffold protein RSP3 were hypophosphorylated (arrow), especially NDK5_{H121A} (triangle). IC140, a subunit of the inner dynein arm, was the loading control. *Con., a WT strain transformed with pNDK5. Dashed line, the arbitrary line to distinguish exogenous proteins expressed from the transgenes. (E) The ratio of rapidly migrating NDK5 from the transgenes relative to total NDK5 in D. Ratio of exogenous protein in Con. strain represents the background. Exo., Exogenous.

NDK5 promotes phosphorylation-related assembly

Absence of entire spokes in some areas in short *ndk5* flagella (Figures 5E and 3D) and hypophosphorylated RSs of reduced abundance in short DN flagella (Figure 8) revealed additional roles of NDK5. First, NDK5 promotes RS assembly, which correlates with protein phosphorylation. Phosphorylation of RSP3 has been implicated in the docking of RSs to axonemes (Gupta *et al.*, 2012). While a major fraction of NDK5 and RSP3 in axonemes is phosphorylated (Figure 3B; Piperno *et al.*, 1981; Gupta *et al.*, 2012), both are largely hypophosphorylated in the cell body (Figure 9; Qin *et al.*, 2004; Gupta *et al.*, 2012). Importantly, hypophosphorylation of both molecules in DN flagella (Figure 8D) shows that NDK5 perturbation results in RSP3 hypophosphorylation and evidently reduces RS assembly.

Second, NDK5 also promotes full-length flagella. *Chlamydomonas ndk5* (Figure 3D) and *ndk5* mouse sperm (Vogel *et al.*, 2012) have short flagella, which can be rescued by the *NDK5* gene (Figure 6). Flagellar length is a multigenic trait ultimately determined by the balance between the rates of assembly and disassembly (Marshall and Rosenbaum, 2001). Flagella defective in one axonemal complex, including absence of the spokehead, are slightly shorter, presumably due to decreased microtubule stability (Kubo *et al.*, 2015). However, *ndk5* flagella are deficient in only three subunits at a similar region and are far shorter than the flagella lacking the entire spokehead. Because NDK5 and RSP3 had identical distributions (Figure 6E; Munier *et al.*, 2003), the simplest explanation is that NDK5 in the RS participates in a distinct length-promotion mechanism. Alternatively, *ndk5*'s length phenotype is still due to a sensitive background in conjunction with the propensity of reduced RS abundance (Figures 5E and 7). Alternatively, NDK5 deficiency reduces the affinity of axonemal proteins and intraflagellar transport (IFT) trains and thus decreases the assembly rate.

We are partial to the possibility that NDK5 promotes axoneme stability or cargo–IFT train affinity via protein phosphorylation. Genetic studies have revealed a number of length-regulating protein kinases and phosphatases (Tam *et al.*, 2013; reviewed by



FIGURE 9: Accumulated hypophosphorylated NDK5 polypeptides in the cell body of the DN strain. (A) A representative Western blot of cell body extracts probed with anti-NDK5₁₋₂₀₁ serum. NDK5 in the DN strain P1 (lane 7) was more abundant than that in the control (lane 5) but similar to that in pf14 (x in lane 8), a control for reduced RS abundance in flagella. Negative control was ndk5 (lane 6). NDK5 bands in axonemes, by themselves or added to cell body extracts (lanes 1-4 and 9), served as markers for phosphorylated NDK5 and hypophosphorylated NDK5 that are endogenous (dot), compared with that from the transgene (triangle) or in pf14 (x). The axoneme sample loaded in lane 3 was one-sixth of the sample loaded in lane 1. (B) A Western blot probed with affinity-purified (Aff. Puri.) anti-NDK5₈₋₅₈₆ polyclonal antibody independently confirmed the hypophosphorylated state of endogenous NDK5 (dot) in the control. The band was undetectable in ndk5. The protein load was shown by Ponceau S stain (bottom).

Wilson et al., 2008; Cao et al., 2009). Many other proteins near the RS base are phosphorylated (Lin et al., 2011). NDK5 may promote phosphorylation of multiple axonemal proteins during the flagellar assembly process, which in turn increases intermolecular affinity, leading to a higher IFT loading rate or greater axoneme stability, and thereby a faster assembly rate (Figure 3, D–F) and full-length flagella. The exacerbation of length phenotypes in the stationary phase (Figures 3, 7, and 8) may reflect declining assembly rates as nutrients are limited, while NDK5 is used to maintain full-length flagella regardless of nutrient availability for algae or sperm.

It is not clear how NDK5 might promote protein phosphorylation. This ability does not require the canonical phosphotransfer pathway of NDK, because *ndk5* is fully rescued by NDK5_{H121A} incapable of the His-dependent phosphotransfer (Figure 6) or even by NDK5_{H121A} lacking one of the additional four residues commonly involved in NTP catalysis. One possible explanation is that NDK5, or perhaps group II NDKs, evolved a distinctive phosphotransfer mechanism while maintaining the canonical one. This is supported by autophosphorylation of the mutated NDK whose conserved His is replaced (MacDonald *et al.*, 1993); distinct residues critical for phosphotransfer in diverse NDKs (Tiwari *et al.*, 2004); and sequence divergence of group I and group II NDKs (Desvignes *et al.*, 2009). Alternatively, NDK5 recruits or activates a protein kinase. In any case, in *ndk5* that lacks most NDK5, a surrogate kinase may phosphorylate RSP3 molecules, albeit less efficiently than NDK5 (Figures 3A, 6C, and 8D). This deficiency may become exacerbated as cultures are reaching stationary phase, leading to the absence of some RSs (Figure 5E). In the DN flagella, the presence of heterodimeric NDK5 may hinder the surrogacy, leading to largely hypophosphorylated RSP3 and, clearly, fewer RSs assembled (Figure 8D). This possibility is consistent with the association of a MAP kinase with the N-terminal extension unique to mammalian RSP3 (Jivan *et al.*, 2009).

The most intriguing observation here is that, although ND-K5_{H121A} is fully functional in ndk5 cells, it elicits a DN effect in WT cells, perturbing phosphorylation of NDK5 and RSP3 and assembly of RSs and flagella. Likewise, $\mathsf{NDK}_{\mathsf{kpn}}$ that has DN effects in Drosophila is functional. The similar abundance of hypophosphorylated NDK5 polypeptides in the cell body of DN strains and pf14 (Figure 9) rules out protein degradation that is attributed to the DN effect of NDK_{kpn} (Biggs et al., 1988; Lascu et al., 1992; Karlsson et al., 1996) or other common DN mechanisms (reviewed by Wilkie, 1994), such as overexpression of inactive molecules. Rather, we propose that NDK5-with or without His-operates with distinct molecular movements, but regardless, the two protomers must act in concert, and thus heterodimers with two functional but incompatible variants exhibit impaired activity. Uncoordinated heterodimersnot limited to phosphorylation—are consistent with NDK mutations in cancer cells (e.g., Chang et al., 1996) and NDK degradation in Drosophila embryos. It is likely that NDK5 heterodimers are unable or less efficient in promoting phosphorylation-related trafficking and assembly. Consistent with this, only a small fraction of NDK5_{H121A} is present in P/S flagella (Figure 8, D and E). Varied rates of P strains among repeated transformations suggest that homo- and heterodimerization of NDK5 is not random.

In fact, intersubunit cross-talk between NDK monomers has been demonstrated (Tepper et al., 1994; Dar and Chakraborti, 2010) and is consistent with classic enzymology analysis that estimated only half of protomers in one NDK oligomer bind to phosphate at one time (reviewed by Parks and Agarwal, 1973). However, this notion has not yet gained the attention it deserves, even though it does not contradict the well-established ping-pong mechanism that depicts phosphotransfer occurring within individual monomers. It is time to reconsider intersubunit cross-talk as a common mechanism of oligomeric NDKs in His-dependent and His-independent actions that remain elusive.

In conclusion, NDK5 is crucial for the assembly of functional motile cilia and flagella, but not because of its canonical NDK activity, which could be redundant. It promotes proper assembly of the RS and full-length flagella as a structural protein and via phosphorylation-related mechanisms. These findings expand the versatility of this ancient molecular platform and provide new clues as to how NDK isoforms are involved in a wide array of cellular processes and how they become impaired.

MATERIALS AND METHODS

Chlamydomonas strains, culture conditions, and biochemistry

WT (cc124 and cc125) and *pf14* allelic mutants—cc613, cc1032, and cc2496—were acquired from the *Chlamydomonas* Resource

Center. The *ndk5* insertional mutant (LMJ.SG0182.001362) and the parental strain (CMJ030) (www.chlamylibrary.org/) were provided by the Jonikas laboratory at the Carnegie Institute of Science (Li *et al.*, 2016). Cells were grown in minimal M media or TAP media (Harris, 2009) as indicated. The light/dark cycle, preparation of flagella, and Western blot analysis were as described previously (Sivadas *et al.*, 2012), except for the addition of protease inhibitor cocktail tablets (cOmplete Mini, EDTA-free; Roche, Basel, Switzerland).

Molecular biology

Characterization of the insertional site in ndk5. PCR was conducted to amplify the PMM-resistance cassette–genome junctions, the upstream *SPL4* gene, and the *NDK5* gene. Templates were genomic DNA freshly prepared from *ndk5* and its parental strain CMJ030. Briefly, cells (~1 μ l) from TAP plates were resuspended in 20 μ l 10 mM EDTA and boiled for 5 min. Following a 30-s vortex and 3-min centrifugation, the supernatant was used in PCR. The primer pairs were as follows:

P1S (GCTCGTGGAGCTCTGAATCT) and P1As (GATACAC-GAACTCCTGCGC) (Li *et al.*, 2016);

P2S (GCTGCATGTAGGAGGCGCCACCTACC) and P2As (GGTG-GCATTCTGGGTGCCGTCCGTTCC); and

P3S (GGCAAATGGAGCTGTTTGGA) and P3As (CCTTAAAGC-GTGCTCCTGAC).

Engineering of cDNA constructs. Two consecutive rounds of PCR were performed using the pMAL-c2 vector (Patel-King *et al.*, 2004) that harbors a partial cDNA encoding aa 8–586 of NDK5 as a template to create a full-length cDNA construct. Two sense primers and one antisense primer are as follows:

NDKS (GCTAGAAAAGACTTTCGCGTTAATCAAGCCAGATGC), NdeS (TC<u>CATATG</u>GCAGAGCTAGAAAAGACTTTCG), and KpnAS (GCGCATCACCGCCACCTCCTTCCG).

The final 1101-base pair NdeS-KpnAS PCR product and the 600-base pair *KpnI-Hind*III 3' fragment released from the pMAL-c2 plasmid were cloned into the pET28 vector between the *Ndel* and *Hind*III sites. This pET28-NDK5 plasmid was converted into pCRS23.2 to express soluble MBP-tagged recombinant NDK5₁₋₂₀₁ in bacteria (Donnelly *et al.* 2006). For expression in *Chlamydomonas*, the coding sequence for NDK5 and the His tag was PCR amplified, and the DNA fragment was used to replace the coding sequence of the LC8 genomic construct that also harbored a PMM-resistance cassette (Yang *et al.*, 2009). The H121 codon in the *NDK5* minigene was mutated to an alanine codon using site-directed mutagenesis and the following primer pair:

NDKH2AS (GGCACCCAGAATGCCACCGCGGGCAGCGACTC-GCC) and NDKH2AAS (GTAGGCGAGTCGCTGCCCGCGGTGGCATTCT-GGGTGCC).

Engineering of genomic constructs. For generation of genomic DNA constructs, an *Ndel-Ndel* fragment containing the *NDK5* gene was released from the 21122 BAC clone (Clemson University Genome Institute) and ligated into the pGEM-T easy vector. The plasmid was further digested with *Eco*RV and *Smal*, and blunt end-ligated to eliminate most of the neighboring *SLP4* gene. The resulting pNDK5 plasmid was converted into pNDK5_{H121A} using site-directed mutagenesis and a primer pair:

NDK5MutS (CCCAGAATGCCACCGCGGGCAGCGACTCGCC-TATCAGC) and

NDK5MutAs (GCTGCCCGCGGTGGCATTCTGGGTGCCGTCC-GTTCC).

An HYG-resistance cassette was PCR-amplified using pHyg3 as a template (Berthold et al., 2002) and inserted into the *Nde*I site in both genomic constructs. The final constructs were 11.3 kb. The primers were as follows:

NdeHygS (GCATATGGATTACGAATTCGATATCAAGCTTCTTTCTTGC) and

NdeHygAs (G<u>CATATG</u>CGCTTCAAATACGCCCAGC).

To express fluorescent RSPs in *Chlamydomonas*, mNeonGreen (Shaner *et al.*, 2013) codons modified in accordance with the *Chlamydomonas* codon bias (Harris *et al.*, 2016) were PCR amplified and cloned into the *Xhol* site inserted before the stop codons of the *NDK5* gene and *RSP3* gene (Sivadas *et al.*, 2012).

Purification of bacterial recombinant proteins

For His-luciferase production, the firefly luciferase gene was amplified from pTRE2-Luc (Clontech, Mountainview, CA) and cloned into the pET28 vector. The plasmids for Hi1-luciferase and His-HSP40 (Yang *et al.*, 2005) were transformed into BL21 (DE3) for protein expression. MBP-His-NDK5₁₋₂₀₁ was expressed as previously described (Donnelly *et al.*, 2006). All His-tagged recombinant proteins were purified with Ni-NTA matrix following the manufacturer's instruction (Qiagen, Hilden, Germany).

Chlamydomonas experiments

Backcross. ndk5 mt(-) was crossed with cc620 or cc125 mt(+) as indicated, using the synchronous gametogenesis method (Snell, 1976).

Transformation. All transformations were conducted using a previously described glass-bead method (Yang *et al.*, 2006) with minor modifications. *NDK5* minigenes at a concentration of 2–8 μ g were transformed into WT cc124 cells, and the transformatis were selected on TAP plates containing 10 μ g/ml PMM. *NDK5* genomic constructs at a concentration of 1–2 μ g were transformed into *ndk5* or its parental strain CMJ030, and the transformants were selected on TAP plates containing 10 μ g/ml HYG. *RSP3* genomic constructs were transformed into *pf14* cells similarly but the transformants were resuspended in 200 μ l double-distilled water in 96-well plates for motility analysis under a stereomicroscope.

Flagellar-length quantification. Cells fixed with 2.5% glutaraldehyde were placed on poly-L-lysine–coated slides for imaging. The lengths of 50 randomly selected flagella were measured using ImageJ software. For flagellar regeneration experiments, pH shock with acetate was used to excise flagella of log-phase cells from TAP medium cultures. After confirmation of deflagellation microscopically, an aliquot of the cells was fixed with 2.5% glutaraldehyde every 15 min for subsequent imaging. The lengths of 20 randomly chosen flagella were measured at each time point.

Light microscopy

Bright-field and fluorescent images were captured with a Nikon Eclipse E600W compound microscope at 400× magnification using a CoolSNAP-ES charge-coupled device camera (Photometrics, Tucson, AZ) and a MetaMorph imaging system (Molecular Devices, Sunnyvale, CA).

Quick-plunge freezing and electron microscopy

Isolated axonemes were quickly frozen in liquid ethane at liquid nitrogen temperatures with the help of a vitrification device (Cryoplunge 3, Gatan, Pleasanton, CA). Holey carbon grids (200 mesh, R3.5/1; Quantifoil Micro Tools GmbH, Großlöbichau, Germany) were used. Gold colloid particles (10 nm) were applied to the sample before freezing as fiducial markers for tomographic reconstruction. Data collection was performed using a JEM2200FS transmission electron microscope (JEOL, Tokyo, Japan) equipped with an in-column energy filter and a field-emission gun. Micrographs were recorded with a 4K × 4K CMOS camera (F416 from TVIPS, Gauting, Germany). Tomographic image series were collected using a previously described procedure (Pigino et al., 2011), except for the use of SerialEM software (Mastronarde, 2005). Image analysis was performed as described previously (Pigino et al., 2011; Bui and Ishikawa, 2013). Initially, 239 particles (subvolumes) were picked from reconstructed tomograms, aligned, and averaged based on 24-nm and 96-nm periodic units. Final average was obtained from 184 bestselected particles. Individual particles included in the final average were used for calculating the 3D variance map using SPIDER software (Frank et al., 1996). Each of the represented individual MTD subaverages contains 12 aligned particles following 96-nm periodic units.

NDK activity assay

The assay measured luminescence emitted from luciferin and was catalyzed by luciferase using ATP generated by NDKs that transferred the γ -phosphate from GTP to ADP (Karamohamed *et al.*, 1999). An aliquot of 60 ng purified recombinant proteins or 1 µg axonemes was added to the 0.2 ml assay buffer (0.1 M Tris-acetate, pH 7.8, 10 mM MgSO₄, 2 mM EDTA, 1 mM dithiothreitol, 1 mg/ml bovine serum albumin, 0.1 mg/ml p-luciferin, 10 ng/ml Ni-NTA-purified His-firefly luciferase, 0.5 mM ADP, and 0.5 mM GTP in a cuvette placed in a Monolight 2010 luminometer [Analytical Luminescence Laboratory, Ann Arbor, MI]). The change in light emission was measured for 10 s and expressed as RLU (light emission/s), as recommended by the manufacturer. Each sample was analyzed in triplicate. The RLU of axonemes for each experimental group was then normalized to that of the WT control and denoted as relative kinase activity (%).

Antibodies

NDK5 Western blots were probed with the rabbit polyclonal antibody raised against Ni-NTA-purified His-tagged NDK5₁₋₂₀₁ unless indicated otherwise. The affinity-purified rabbit polyclonal antibodies for NDK5₈₋₅₈₆ and the antibodies for the other proteins were described in previous studies (LeDizet and Piperno, 1995; Patel-King et al., 2004; Yang et al., 2006). The 3G3 was a mouse mAb raised against 120-kDa spoke proteins spot-purified from two-dimensional gels of axonemes (Williams et al., 1989). This study showed that 3G3 recognized NDK5. Anti-IC1 was a mouse mAb (King et al., 1991).

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