

Structure and Function of Human DnaJ Homologue Subfamily A Member 1 (DNAJA1) and Its Relationship to Pancreatic Cancer

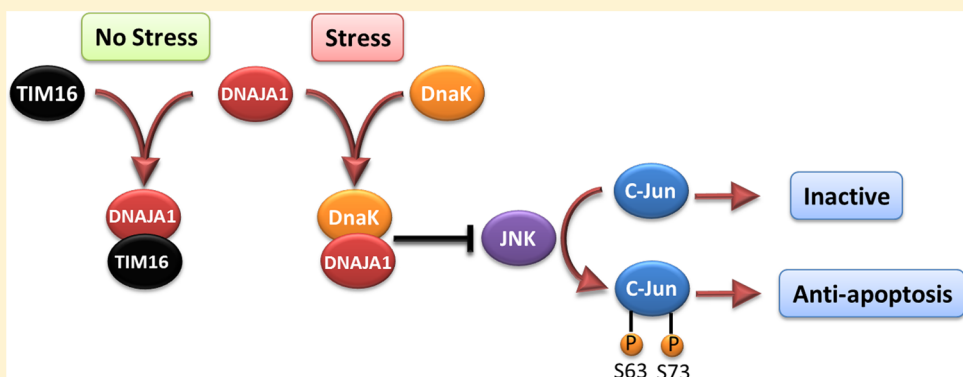
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ABSTRACT: Pancreatic cancer has a dismal 5 year survival rate of 5.5% that has not been improved over the past 25 years despite an enormous amount of effort. Thus, there is an urgent need to identify truly novel yet druggable protein targets for drug discovery. The human protein DnaJ homologue subfamily A member 1 (DNAJA1) was previously shown to be downregulated 5-fold in pancreatic cancer cells and has been targeted as a biomarker for pancreatic cancer, but little is known about the specific biological function for DNAJA1 or the other members of the DnaJ family encoded in the human genome. Our results suggest the overexpression of DNAJA1 suppresses the stress response capabilities of the oncogenic transcription factor, c-Jun, and results in the diminution of cell survival. DNAJA1 likely activates a DnaK protein by forming a complex that suppresses the JNK pathway, the hyperphosphorylation of c-Jun, and the anti-apoptosis state found in pancreatic cancer cells. A high-quality nuclear magnetic resonance solution structure of the J-domain of DNAJA1 combined with a bioinformatics analysis and a ligand affinity screen identifies a potential DnaK binding site, which is also predicted to overlap with an inhibitory binding site, suggesting DNAJA1 activity is highly regulated.

Despite the decline of cancer-related mortality in the past decade, effective approaches to early diagnosis and treatment of pancreatic cancer remain elusive. Although it accounts for only 3% (43000 new cases every year) of all cancers, pancreatic cancer is the fourth leading cause of cancer death in the United States (37000 deaths annually) and has the highest mortality rate of any cancer.^{1,2} Those with an operable early stage of the disease have a 5 year survival rate of ~20%.^{2,3} Unfortunately, 80% of all pancreatic cancer diagnoses indicate an advanced stage of the disease that is beyond the point of surgery.^{1,3,4} Inoperable forms of pancreatic cancer have a 5 year survival rate of only 3%. The difficulty in detecting or diagnosing pancreatic cancer has several causes: the early stages of pancreatic cancer do not typically exhibit symptoms;

the symptoms that do occur are often similar to those of other illnesses; and the location of the pancreas behind other organs can hinder detection.¹

Most patients with advanced pancreatic cancer are treated with chemotherapy based on gemcitabine, which is a cytotoxic nucleoside drug that primarily inhibits DNA synthesis.⁵ However, this treatment is only mildly effective for patients with an advanced stage of pancreatic cancer and provides an only 5.91 month increase in the median survival rate.⁶ Also,

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gemcitabine-resistant forms of pancreatic cancer and acquired resistance during treatment are common problems.⁷ Correspondingly, there have been numerous attempts to combine gemcitabine with other cytotoxic agents, such as 5-fluorouracil or capecitabine. However, these approaches have been mostly unsuccessful.⁸ It is apparent that a cytotoxic approach to treating pancreatic cancer is not an effective therapy. Therefore, identifying novel, but druggable, protein targets for the treatment of pancreatic cancer and improving the quality of life for patients are essential needs.

The DnaJ proteins, also known as heat shock protein 40 (Hsp40 or Hsc40), are proteins originally identified in *Escherichia coli* that act as cochaperones to the molecular chaperone DnaK (Hsp70), which is responsible for several cellular processes such as rescuing misfolded proteins, folding polypeptide chains, transport of polypeptides through membranes, assembly and disassembly of protein complexes, and control of regulatory proteins.^{9–11} DnaJ primarily facilitates the hydrolysis of ATP from DnaK that is necessary for the chaperone activity of DnaK.^{11–13} In general, J-domain proteins modulate protein assembly, disassembly, and translocation.¹⁴

Human protein DnaJ subfamily A member 1 (DNAJA1) has been shown to associate on its own with unfolded polypeptide chains and prevent their aggregation,¹⁵ to regulate androgen receptor signaling and spermatogenesis in mice,¹⁶ and to contribute to the resistance of glioblastomas to radiotherapy.¹⁷ DNAJA1 has also been targeted as a biomarker for pancreatic cancer to evaluate the effects of farnesyl protein transferase inhibitors^{18,19} and has been shown to be downregulated 5-fold in a genomics analysis of pancreatic cancer cells relative to normal healthy cells and cells undergoing pancreatitis.²⁰ Additionally, DNAJA1 appears to be involved in importing proteins into the mitochondria.^{21,22} Of note, the mitochondrial pathway to apoptosis protects against cancer and requires importing apoptotic factors into the mitochondrial membrane.^{23–26} Additionally, DNAJA1 is an interesting target based on its association with DnaK, which is expressed abundantly in various tumors and can potentially promote tumorigenesis by inhibiting cell death.^{27–32} However, there have been no studies of whether the conserved J-domain of DnaJ alone has any role in cancer biology independent of DnaK.¹⁰

Thus, DNAJA1 was selected for further analysis as a potentially interesting therapeutic target for the treatment of pancreatic cancer. The potential importance of DNAJA1 to pancreatic cancer was demonstrated by stress response cell-based assays using cell lines overexpressing DNAJA1. DNAJA1 was shown to regulate the hyperphosphorylation of c-Jun and cell survivability under stress. The resulting nuclear magnetic resonance (NMR) structure of the J-domain of DNAJA1 (DNAJA1-JD) combined with a detailed bioinformatics analysis predicted the location of overlapping activating and inhibitory protein binding sites. A subsequent NMR-ligand affinity screen provided further experimental support for the existence of the predicted inhibitory protein binding sites and also demonstrated that the J-domain of DNAJA1 is a potential druggable target. In total, our results suggest DNAJA1 activity is highly regulated and that DNAJA1 is involved in a stress response signaling pathway that suppresses the anti-apoptosis state found in pancreatic cancer cells.

EXPERIMENTAL PROCEDURES

Effect of DNAJA1 Overexpression on Cellular Stress Modulation in Pancreatic Cancer Cells.

MiaPaCa2 cells

were obtained from the American Type Culture Collection (Rockville, MD). Cells were cultured as previously described.³³ Briefly, MiaPaCa2 cells were maintained in Dulbecco's modified Eagle's medium (Life Technologies, Carlsbad, CA) supplemented with 10% heat-inactivated fetal bovine serum (FBS), nonessential amino acids, sodium pyruvate, and penicillin/streptomycin in a 37 °C incubator with 5% CO₂. To stably express the full-length His-tagged DNAJA1 construct, retroviral transductions were conducted essentially as described previously.³⁴ The DNAJA1 construct was purchased from the Arizona State University Biodesign Institute (<http://dnasu.org/DNASU/Home.jsp>).

Cell lysates were prepared by scraping cells (80–90% confluent) into lysis buffer [50 mM Tris-HCl (pH 7.5), 0.15 M NaCl, 1% (v/v) Triton X-100, 1% (w/v) sodium deoxycholate, 0.1% (w/v) SDS, 5 mM EDTA, and 1 mM phenylmethanesulfonyl fluoride]. Lysates were incubated, on ice, for 30 min and centrifuged at 4 °C for 15 min at 13000 rpm to remove cell debris. Supernatants were transferred to fresh tubes, and the protein content was determined using the Bradford protein assay reagent (Bio-Rad, Hercules, CA) with various concentrations of bovine serum albumin as standards. Cell lysates were stored at –20 or –80 °C. Cell lysate proteins were resolved on 10 or 12% Tris-glycine denaturing polyacrylamide gels in 1× sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) buffer (1 g/L SDS, 3 g/L Tris base, and 14.4 g/L glycine). Western blotting was performed as previously described.^{35,36} The membranes were probed with primary antibodies against phosphoS63-c-Jun [2361 (Cell Signaling Technology, Danvers, MA)], total c-Jun [9165 (Cell Signaling Technology)], the His tag [2365 (Cell Signaling Technology)], and tubulin [E7 (Developmental Studies Hybridoma Bank, Iowa City, IA)]. The primary antibody treatments were performed overnight at 4 °C at 1:1000 dilutions.

Cell survival was evaluated by the MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] assay as described elsewhere.³⁷ Cells were plated in triplicate in 96-well plates at a density of 5000 cells/well and incubated at 37 °C. Twelve hours later, cells were treated with anisomycin (10 µg/mL) or solvent control [dimethyl sulfoxide (DMSO)] and incubated for 24 h. At the end point, the culture medium was removed and 20 µL of the MTT solution [5 mg/mL (Sigma-Aldrich, St. Louis, MO)] was added per well, followed by a 2 h incubation. MTT was removed, and 200 µL of DMSO was added to each well to dissolve formazan. The formazan optical density was determined by utilizing a microplate reader (FLUOstar Omega, BMG LABTECH, Cary, NC) at a wavelength of 540 nm.

Solution Structure of the DNAJA1 J-Domain. The full-length human DNAJA1 protein (397 amino acids) has been targeted by the Northeast Structural Genomics Consortium [NESG (<http://www.nesg.org>)] for structural elucidation as target HR3099 (UniProt entry P31689) (Figure 1).

Samples of approximately 100% ¹⁵N-enriched, 5% ¹³C-enriched, and uniformly ¹³C- and ¹⁵N-enriched DNAJA1-JD [77 amino acids with 10 non-native residues, MGHHHHHH-SH, at the N-terminus for purification (NESG entry HR3099K)]^{38,39} were produced using the standard protocols of the NESG.^{38,39} The protein construct containing the sequence for DNAJA1-JD was transformed into BL21(DE3)+Magic cells. The soluble fraction of the lysed cells was collected and purified with a Ni⁺ affinity column (HisTrap HP 5 mL column, GE Healthcare, Pittsburgh, PA) and a gel filtration column

MVKETTYDVLGVKPATQEELKKAYRKLAALKYHPDKNPNEGEKFQKIS
QAYEVLSDAKKRELYDKGGQAIKEGGAGGGFGSPMDIFDMFFGGGGRM
QRERRGKNNVHVQLSNTLEDLYNGATRKLALQKNVICDKCEGRGGKKGAV
ECCPNCRGTGMQIRIHQIGPGMVQQIQSVCMECQGHGERISPKDRCKSC
NGRKIVREKKILEVHIDKGMDQKITFHGEGLDQEPLGEPGDIIVLHQ
KDHAFTTRRGEDLFMCMIDIQLVEALCGFQKPISTLDNRTIVTSHPGQI
VKHGDIKCVLNEGMPYRRPYEKGRLLIEFKVNFPENGFLSPDKLSSLLE
KLLPERKEVEETDEMDQVELVDFDPNQERRRHYNGEAYEDDEHHPRGGV
QCQTS

Figure 1. Protein sequence of DNAJA1 (UniProt entry P31689). The red residues are the 67 amino acids of the J-domain (DNAJA1-JD).

(HiLoad 26/60 Superdex 75 prep grade, GE Healthcare). The sample identity and isotopic enrichment were validated by matrix-assisted laser desorption ionization time-of-flight mass spectrometry, and the homogeneity (>97%) was assessed by SDS-PAGE. The NMR protein sample was stored in a sealed Shigemi tube with 20 mM 2-(4-morpholino)ethanesulfonic acid [MES (Sigma-Aldrich)] buffer [pH 6.5 (uncorrected)] with 10% D₂O (Sigma-Aldrich), 0.02% NaN₃ (Sigma-Aldrich), 10 mM dithiothreitol [DTT (Sigma-Aldrich)], 5 mM CaCl₂ (Sigma-Aldrich), 100 mM NaCl (Sigma-Aldrich), and 50 μM 4,4-dimethyl-4-silapentane-1-sulfonic acid [DSS (Sigma-Aldrich)].

NMR experiments used for the protein backbone and side-chain assignments of DNAJA1-JD were collected at 298 K on a 600 MHz Bruker Avance spectrometer equipped with a 5 mm TXI probe. The backbone and side-chain assignments were completed using the standard triple-resonance approach^{40–42} of the following NMR experiments: two-dimensional (2D) ¹H-¹⁵N HSQC, 2D ¹H-¹³C HSQC, and three-dimensional (3D) HNCO; HN(CA)CO; HNCA; HN(CO)CA; CBCA(CO)NH; CBCANH; HNHA; HBHA(CO)NH; CC(CO)-NH; HCC(CO)NH; HCCH-COSY; and HCCH-TOCSY. ¹⁵N-edited NOESY-HSQC and ¹³C-edited NOESY-HSQC experiments (mixing times of 120 ms) were conducted on a 500 MHz Bruker Avance DRX spectrometer equipped with a triple-resonance, Z-axis gradient cryoprobe to identify nuclear Overhauser effects (NOEs). Amide hydrogen exchange rates were also evaluated on the 500 MHz Bruker Avance DRX spectrometer using the (CLEANEX-PM)-FHSQC experiment.

The NMR experimental data were processed using NMRPipe⁴³ and evaluated in CCPNMR Analysis (<http://www.ccpn.ac.uk>).⁴⁴ An initial model of DNAJA1-JD was generated from the protein backbone resonances using CS ROSETTA on the WeNMR GRID-enabled web portal (<http://www.enmr.eu/webportal>).^{45–47}

The initial model of DNAJA1-JD was refined with XPLOR-NIH version 2.31⁴⁸ using the following experimental restraints: 1070 NOE distance restraints, 50 H-bond distance and angle restraints, 38 ³J_{NHα} coupling constant restraints, 127 ¹³Ca/¹³Cβ chemical shift restraints, and 116 dihedral angle restraints predicted from TALOS+.⁴⁹ All peptide bonds were constrained to be planar and *trans*. A total of 400 structures were calculated, with the 20 lowest-energy structures being subjected to explicit water refinement based on the RECOORD protocols.⁵⁰ An average DNAJA1-JD structure was calculated on the basis of the average atom coordinates of the 20 water-refined structures and subsequently minimized using the same explicit water refinement described above.

The resulting structures were evaluated using the RPF webserver (<http://nmr.cabm.rutgers.edu/rpf/>)⁵¹ and PSVS software suite (http://psvs-1_5-dev.nesg.org/),⁵² which in-

cludes Verify3D,⁵³ ProsaII,⁵⁴ PROCHECK,⁵⁵ and Molprobity.⁵⁶ The three-dimensional structures of the proteins are represented here using the UCSF Chimera package from the Resource for Biocomputing, Visualization, and Informatics at the University of California, San Francisco (<http://www.cgl.ucsf.edu/chimera/>).⁵⁷ ClustalW⁵⁸ was used (with default settings) to align the sequences of DNAJA1-JD with the J-domains of four homologous proteins: *E. coli* DnaJ J-domain [Protein Data Bank (PDB) entry 1XBL],⁵⁹ *Homo sapiens* Hsp40 (HDJ-1) J-domain (PDB entry 1HDJ),⁶⁰ *H. sapiens* HSJ1a (PDB entry 2LGW),⁶¹ and *H. sapiens* DnaJ subfamily C member 12 (PDB entry 2CTQ). Electrostatic surface potentials of the protein were calculated using Delphi.⁶² The identification of evolutionarily conserved residues using both sequence and structure was performed using the ConSurf server with default settings.⁶³

Identification of a Ligand Binding Site on the DNAJA1 J-Domain.

To experimentally determine a potential functional epitope or functionally relevant ligand binding site, we used a high-throughput NMR ligand affinity screen using a function-based compound library.^{64,65} The one-dimensional (1D) ¹H line broadening ligand-based screen and 2D ¹H-¹⁵N HSQC protein-based screen follow the same procedure outlined elsewhere^{66,67} using 10 and 30 μM DNAJA1-JD (5% ¹³C-labeled and 100% ¹⁵N-labeled), respectively. The concentrations for the compounds were 100 μM in the 1D ¹H line broadening screen and 400 μM in the 2D ¹H-¹⁵N HSQC screen.

The NMR spectra for the ligand affinity screens were recorded on a 500 MHz Bruker Avance DRX spectrometer equipped with a triple-resonance, Z-axis gradient cryoprobe with a Bruker autotune and match (ATM) and BACS-120 sample changer. All 1D ¹H NMR spectra were processed with the ACD/NMR Processor (ACD/Laboratories), and 2D ¹H-¹⁵N HSQC spectra were processed with NMRPipe⁴³ and visualized in CCPNMR Analysis.⁴⁴

Chemical shift perturbations (CSPs) between the free and ligand-bound 2D ¹H-¹⁵N HSQC spectra of DNAJA1-JD were used to define the consensus binding site. The CSPs were calculated using a common weighting approach⁶⁸ based on changes in ¹⁵N and ¹H chemical shifts that occur upon ligand binding. AutoDock version 4.2.3^{69–71} with the AutoDockTools version 1.5.4^{71,72} graphical interface was used to calculate 120 protein-ligand costructures, which were filtered using AutoDockFilter⁷³ to identify the costructures that best agree with the experimental CSPs.

RESULTS AND DISCUSSION

Effect of DNAJA1 Overexpression on Cellular Stress Modulation in Pancreatic Cancer Cells. DNAJA1 is related to the cellular response to stress, a common event in the tumor cells due to fluctuations in the nutrient and oxygen levels in the tumor microenvironment or due to therapy, and can govern the response of the tumor to chemo- and radiotherapies. Hence, we performed cell-based functional assays of c-Jun phosphorylation to identify DNAJA1-mediated activation of the stress response pathway in cancer cells. c-Jun is a member of the JNK signaling pathway, and its transcriptional activity and expression are primarily regulated by the phosphorylation of two serines, Ser63 and Ser73.^{74–77} c-Jun regulates a range of cellular processes, including apoptosis, tumorigenesis, and cell proliferation, which includes protecting cells from stress-induced cell death.^{75,77} This makes c-Jun both a positive and negative

regulator of cell death. However, in several cancers, c-Jun has been shown to inhibit apoptosis, leading to the uncontrolled growth typical of cancer.^{78–80}

Human MiaPaCa2 cells stably expressing full-length DNAJA1 or the vector control were subjected to a 1 h incubation with anisomycin (370 nM), a protein synthesis inhibitor, or UV treatment at 20 or 50 J/m² doses to mimic stress. Activation of the stress-induced JNK pathway was measured by evaluating the downstream phosphorylation of c-Jun 20 min after treatment. The results indicate that the overexpression of DNAJA1 (i.e., in cell line MiaPaCa2-DNAJA1) diminishes the level of anisomycin and UV-induced c-Jun phosphorylation at residue Ser63 (Figure 2A). Please

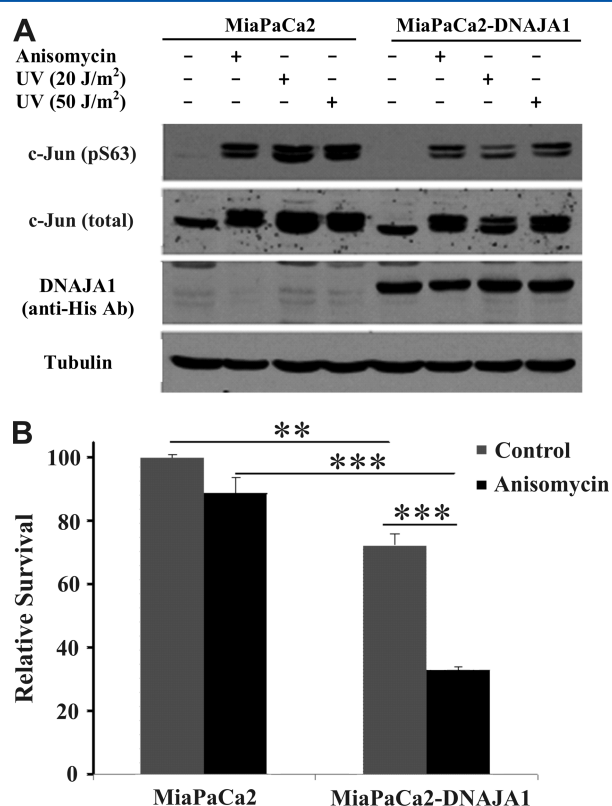


Figure 2. (A) Expression of full-length DNAJA1 (UniProt entry P31689) in MiaPaCa2 pancreatic cancer cells suppresses the activation of c-Jun in response to anisomycin (370 nM) or UV treatment (20 or 50 J/m² doses). Twenty minutes after being treated, cells were subjected to lysis, and the levels of phospho-c-Jun (S63) and total c-Jun were evaluated by Western blotting. Expression levels of exogenously expressed His-tagged DNAJA1 were evaluated by immunoblotting with the anti-His antibody, while immunoblotting with the anti-tubulin antibody was performed as a loading control. (B) Expression of full-length DNAJA1 in MiaPaCa2 pancreatic cancer cells decreases the level of survival in response to anisomycin treatment-induced stress. The level of cell survival was measured by the MTT assay 24 h after treatment (**p < 0.01; ***p < 0.001).

note the doubling of the c-Jun bands in Figure 2A is a result of different phosphorylation states of c-Jun and, correspondingly, different gel migration rates. Cell survival was also evaluated by performing MTT assays 24 h after treatment with anisomycin. The results indicate that DNAJA1 expression decreased the level of cell survival under conditions of anisomycin treatment (Figure 2B).

Expression of the DNAJA1 protein in MiaPaCa2 pancreatic cancer cell lines results in the cells being more susceptible to the stress-induced (UV-induced DNA damage or anisomycin-induced inhibition of protein synthesis) diminution of the level of cell survival (Figure 2). DNAJA1 expression also suppresses the phosphorylation-mediated activation of the oncogenic transcription factor, c-Jun, which is often overexpressed and hyperphosphorylated in cancer^{81–83} and has an essential role in pancreatic cancer.^{84,85}

Interestingly, heat shock proteins have also been shown to be regulators of apoptosis, where DnaK (Hsp70) suppresses JNK activity.^{86–88} Thus, our preliminary results suggest DNAJA1 potentially stimulates the DnaK suppression of a JNK-induced anti-apoptotic signaling pathway by forming a complex with DnaK. This hypothesis is consistent with the 5-fold down-regulation of DNAJA1 in pancreatic cancer cells, the resulting suppression of c-Jun phosphorylation, and the corresponding susceptibility to stress-induced apoptosis by expressing DNAJA1. Obviously, protection from stress-induced apoptosis by downregulating DNAJA1 would be beneficial because cancer cells, by definition, exist in a stressful environment. To further investigate this role of DNAJA1 in pancreatic cancer, we determined an NMR solution structure.

Solution Structure of the DNAJA1 J-Domain. The backbone resonance assignments were determined for 63 (98.4%) of the assignable 64 amino acid residues (excluding the 10 non-native residues to the N-terminal side of the His tag, and the three Pro residues). The one amino acid for which the amide could not be assigned, Met11, is the first residue following the His tag. The side-chain assignments were completed using a combination of CC(CO)NH, HCC(CO)-NH, HCCH-COSY, and HCCH-TOCSY 3D NMR experiments. The backbone and side-chain assignments, not including the 10-residue His tag, were nearly complete with 63 of 67 N, 63 of 64 HN, 67 of 67 C α , 70 of 70 H α , 63 of 64 C β , 113 of 115 C β , 43 of 62 C γ , 66 of 74 H γ , 26 of 48 C δ , 40 of 54 H δ , 11 of 21 C ε , 14 of 34 H ε , 0 of 9 C ζ , and 0 of 1 H ζ atoms assigned.

Using the backbone resonance assignments, a model of DNAJA1-JD was generated using CS-ROSETTA, which utilizes chemical shifts to select protein fragments from the PDB followed by Monte Carlo assembly and relaxation by Rosetta.^{46,47} This tool has been shown to be effective in predicting protein structures for small proteins (\leq 16 kDa). The model generated from CS-ROSETTA exhibited the same secondary structure as most Dnaj proteins and agreed with the secondary structure predicted from TALOS+. All available backbone and side-chain chemical shift assignments have been deposited into the Biological Magnetic Resonance Data Bank [BMRB (<http://www.brbm.wisc.edu>)] as entry 19163.

The solution structure of DNAJA1-JD was calculated using 1120 distance restraints, 116 dihedral restraints, 127 C α /C β chemical shift restraints, and 38 3 J_{NH α} coupling constant restraints. The restraints used during the structure calculation are summarized in Table 1. XPLOR-NIH was used to calculate 400 structures, and the 20 lowest-energy structures were selected for further refinement in water using the RECOORD protocol implemented in XPLOR-NIH. The resulting ensemble and average structures (Figure 3A,B) agreed well with the NMR data, where the experimental restraints had low root-mean-square deviations (rmsds) (Table 1). The water-refined average structure had no NOE violations greater than 0.5 Å or dihedral violations greater than 5°. The water-refined ensemble of 20 structures had a backbone rmsd of 0.709 ± 0.119 Å from

Table 1. Structure Calculation Statistics^a

	$\langle \text{SA} \rangle$	$(\overline{\text{SA}})_r$
rmsd for distance restraints (experimental) (Å)		
all (1120)	0.083 ± 0.001	0.082
inter-residue sequential ($ i - j = 1$) (269)	0.082 ± 0.003	0.086
inter-residue short-range ($1 < i - j < 5$) (221)	0.078 ± 0.005	0.073
inter-residue long-range ($ i - j \geq 5$) (80)	0.108 ± 0.008	0.098
intraresidue (500)	0.084 ± 0.002	0.085
H-bonds (50)	0.038 ± 0.008	0.029
rmsd for dihedral angle restraints (deg) (116)	0.027 ± 0.044	0.00
rmsd for $C\alpha$ and $C\beta$ shifts restraints (ppm) (127)	0.873 ± 0.036	0.907
rmsd for $^3J_{\text{NH}\alpha}$ restraints (Hz) (38)	0.536 ± 0.038	0.540
rmsd (covalent geometry)		
bonds (Å)	0.007 ± 0.000	0.007
angles (deg)	0.649 ± 0.015	0.647
impropers (deg)	0.794 ± 0.037	0.778
energy (kcal/mol)		
total	-2620.08 ± 70.18	-2834.43
bond	37.44 ± 2.35	36.16
angle	94.07 ± 4.69	95.65
dihedral	0.02 ± 0.04	0.00
impropers	33.44 ± 2.77	31.56
van der Waals	-262.40 ± 10.76	-273.46
NOE	230.11 ± 7.09	226.01
$^3J_{\text{NH}\alpha}$	10.99 ± 1.60	11.08
$C\alpha$ and $C\beta$ shifts	49.16 ± 4.30	52.92

^a $\langle \text{SA} \rangle$ represents the final 20 water-refined simulated annealing structures. $(\overline{\text{SA}})_r$ represents the water-refined average structure of all 20 water-refined structures.

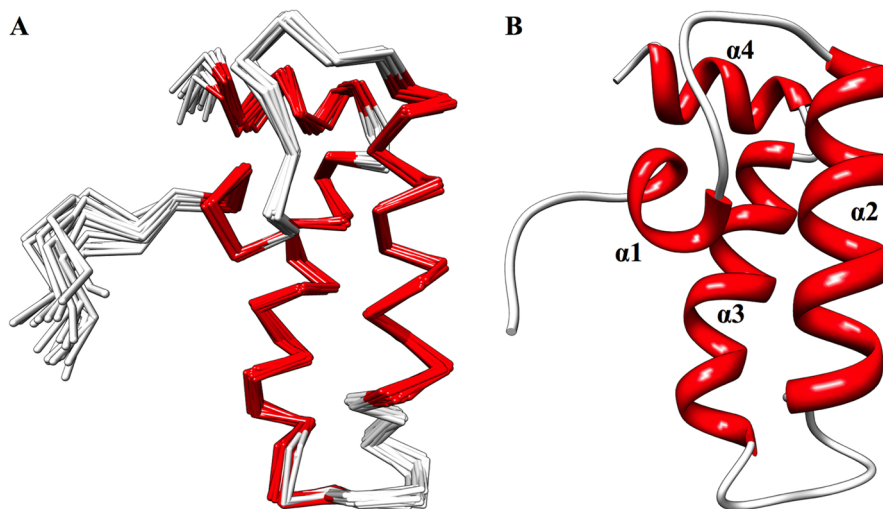


Figure 3. (A) Overlay of the backbone trace of the 20 lowest-energy, water-refined structures. (B) Ribbon representation of the average structure generated from the average atomic coordinates of the 20 lowest-energy, water-refined structures, followed by water refinement of the average structure. Both structures are colored according to secondary structure: red for α -helix and white for loop.

Table 2. Atomic Root-Mean-Square Differences^a

	full protein (residues 11–77)		secondary structure	
	backbone atoms	all heavy atoms	backbone atoms	all heavy atoms
$\langle \text{SA} \rangle$ vs $\overline{\text{SA}}$	0.709 ± 0.119	1.417 ± 0.110	0.399 ± 0.086	1.232 ± 0.134
$\langle \text{SA} \rangle$ vs $(\overline{\text{SA}})_r$	0.888 ± 0.136	1.739 ± 0.139	0.639 ± 0.155	1.605 ± 0.131
$(\overline{\text{SA}})_r$ vs $\overline{\text{SA}}$	0.649	1.217	0.523	1.183

^a $\langle \text{SA} \rangle$ represents the final 20 water-refined simulated annealing structures. $\overline{\text{SA}}$ represents the average structure of all 20 water-refined structures. $(\overline{\text{SA}})_r$ represents the water-refined average structure of all 20 water-refined structures.

the unrefined average coordinates. This result improves to 0.399 ± 0.086 Å when only the residues involved in the more stable secondary structure elements are evaluated, which indicates the consistency of the structure calculation using the experimental restraints (Table 2). Comparing the original CS-ROSETTA homology model to the final water-refined average structure showed backbone rmsds of 1.803 Å (full protein) and 0.889 Å (secondary structure).

The RPF webserver and PSVS software suite were used to verify the quality of the ensemble and average structures (Table 3). Few unreasonable atom clashes were identified by the

Table 3. Structural Evaluation^a

	$\langle SA \rangle$	$\langle \overline{SA} \rangle_r$
PSVS Z-score ^b		
Verify3D	-0.80	-0.16
ProsaII (-ve)	2.40	2.52
Procheck (ϕ and ψ)	1.65	1.46
Procheck (all)	1.48	1.54
MolProbity clash score	-1.59	-1.56
RPF scores ^c		
Recall	0.968 ± 0.001	0.965
Precision	0.762 ± 0.006	0.737
F-measure	0.852 ± 0.004	0.836
DP-score	0.733 ± 0.020	0.656
Ramachandran space ^d (%)		
most favored regions	98.5	98.5
allowed regions	1.5	1.5
disallowed regions	0.0	0.0

^a $\langle SA \rangle$ represents the final 20 water-refined simulated annealing structures. $\langle \overline{SA} \rangle_r$ represents the water-refined average structure of all 20 water-refined structures. ^bCalculated with PSVS (http://psvs-1_5-dev.nesg.org/). More positive scores are better. ^cCalculated with RPF (<http://nmr.cabm.rutgers.edu/rpf/>). A DP-score of >0.75 and an F-measure of >0.8 are consistent with high-quality structures.

^dCalculated with the Molprobity module in PSVS.

Molprobity module, and our DNAJA1-JD NMR structure has a very good Z-score (-1.59) compared to the average Z-score for NMR structures in the PDB (-10.74).⁵² This is actually comparable to the Z-scores of medium-resolution X-ray crystal structures in the PDB (-1.39).⁵² An evaluation of the probable dihedral angles expected for each residue using PROCHECK also shows excellent Z-scores (1.65 for ϕ and ψ dihedrals and 1.48 for all dihedrals). Additionally, 98.5% of the residues have backbone dihedral angles in the most favored regions of Ramachandran space; only one residue, Val12, has backbone ϕ and ψ values in the more generously allowed region. The discriminating power (DP-score), Recall, Precision, and F-measure scores provide global measures of the goodness of fit between the DNAJA1-JD NMR structure and the NOESY peak list, where a DP-score of >0.75 and an F-measure of >0.8 are consistent with high-quality structures. The DP-score and F-measure calculated for the DNAJA1-JD NMR structure are 0.852 and 0.733, respectively. Overall, the results indicate that the final ensemble and average models for DNAJA1-JD are good-quality structures with few to no unreasonable structural features. The coordinates of the water-refined ensemble have been deposited in the PDB as entry 2M6Y.

The secondary structure and fold for DNAJA1-JD are similar to those of other J-domains found in DnaJ homologues. In effect, our DNAJA1-JD structure adopts the characteristic J-

domain found in most species. The structure consists of four α -helices: residues 17–21 ($\alpha 1$), 29–42 ($\alpha 2$), 52–65 ($\alpha 3$), and 68–75 ($\alpha 4$) (Figure 3B). The loop between helices $\alpha 2$ and $\alpha 3$ (residues 43–51) contains the highly conserved His-Pro-Asp (HPD) motif (residues 44–46).

The structures of 28 DnaJ proteins in various organisms have been determined. Most of these proteins (16 structures) are from humans, with the majority belonging to DnaJ subfamily B (six) and DnaJ subfamily C (eight). The tertiary structure of DNAJA1-JD was compared to a few representative structures: *E. coli* DnaJ J-domain (PDB entry 1XBL), human DnaJ homologue subfamily B member 1 J-domain (PDB entry 1HDJ), human DnaJ homologue subfamily B member 2 J-domain (PDB entry 2LGW), and human DnaJ homologue subfamily C member 12 J-domain (PDB entry 2CTQ). All four proteins have essentially the same tertiary structure as DNAJA1-JD with PDBeFold Z-scores of 5.148 (2.08 Å rmsd), 6.872 (1.65 Å rmsd), 4.541 (2.24 Å rmsd), and 6.904 (1.37 Å rmsd), respectively (Figure 4A–D). The different DnaJ homologue subfamilies do not appear to exhibit a significant difference in structure, especially because the best matched structure to DNAJA1 belongs to DnaJ homologue subfamily C. Thus, the structural and functional difference between these subfamilies likely occurs in the other DnaJ domains.

The human DNAJA1-JD sequence is also very well conserved in J-domains from other organisms, with 56 proteins having $\geq 49\%$ identical sequences. The sequences of representative DnaJ proteins 1XBL, 1HDJ, 2LGW, and 2CTQ have 51, 56, 47, and 32% identical sequences, respectively (Figure 4E). As expected, all five proteins have the highly conserved HPD motif. Additionally, helix $\alpha 2$ is highly conserved except for protein 2CTQ. This is interesting and potentially functionally relevant because helix $\alpha 2$ tends to be positively charged and represents a possible binding site for DnaK (Hsp70).

Identification of Potential Protein Binding Sites on the DNAJA1 J-Domain. One of the primary functions for DnaJ is to stimulate the ATPase activity of DnaK. The primary function of the J-domain of DnaJ is to bind to the ATPase domain on DnaK. As previously mentioned, the main feature of J-domain proteins is a highly conserved HPD motif, which may indicate its importance in binding to DnaK. Thus, identifying a ligand binding site that overlaps with this proposed DnaK binding site would provide further validation of the location and existence of a potential DnaK binding site. Critically, a specific DnaK or DnaK-like protein that binds DNAJA1 has not been identified. It is simply hypothesized to exist on the basis of analogy with other DnaJ proteins.¹¹

The PDB contains only one example of a J-domain in complex with DnaK. This example (PDB entry 2QWN) is a crystal structure of the bovine auxilin (DnaJ homologue subfamily C) J-domain chemically cross-linked with bovine DnaK at the conserved HPD motif.^{9,89} A sequence alignment of the residues between the bovine auxilin J-domain and DNAJA1-JD allows for a prediction of the proposed DnaJ–DnaK interaction site (Figure 5A). This proposed binding site includes the highly conserved HPD motif. However, there is some contention about whether the cross-linked complex is biologically relevant or whether auxilin accurately represents most interactions of DnaJ with DnaK.^{89–91}

A previous NMR analysis revealed chemical shift perturbations (CSPs) in 2D ^1H – ^{15}N HSQC experiments when the *E. coli* DnaJ J-domain is bound to *E. coli* DnaK.⁹² The majority of

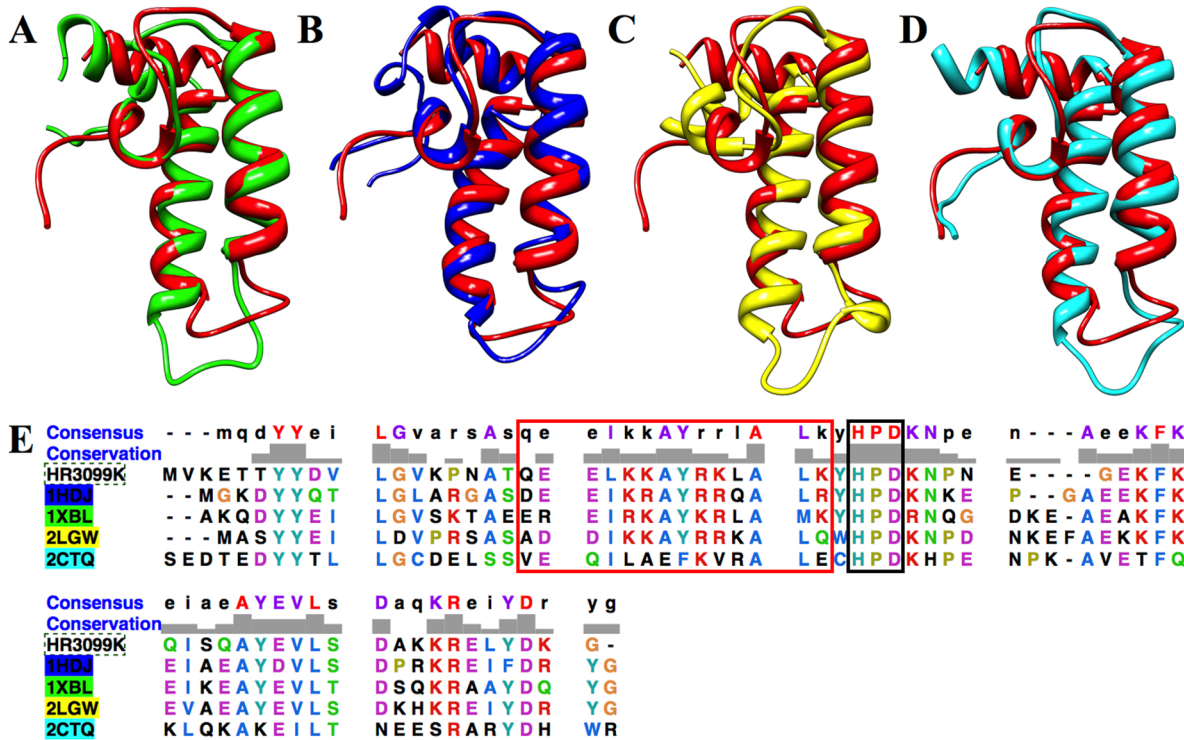


Figure 4. Overlay of the ribbon structure for DNAJA1-JD (red) with (A) the *E. coli* DnaJ J-domain (PDB entry 1XBL), (B) the *H. sapiens* DnaJ homologue subfamily B member 1 J-domain (PDB entry 1HDJ), (C) *H. sapiens* DnaJ homologue subfamily B member 2 (PDB entry 2LGW), and (D) *H. sapiens* DnaJ homologue subfamily C member 12 (PDB entry 2CTQ). (E) ClustalW comparison of DNAJA1-JD (HR3099K) with PDB entries 1HDJ (blue), 1XBL (green), 2LGW (yellow), and 2CTQ (cyan). The highly conserved HPD sequence is outlined with a black box. The residues that make up helix $\alpha 2$ are outlined with a red box.

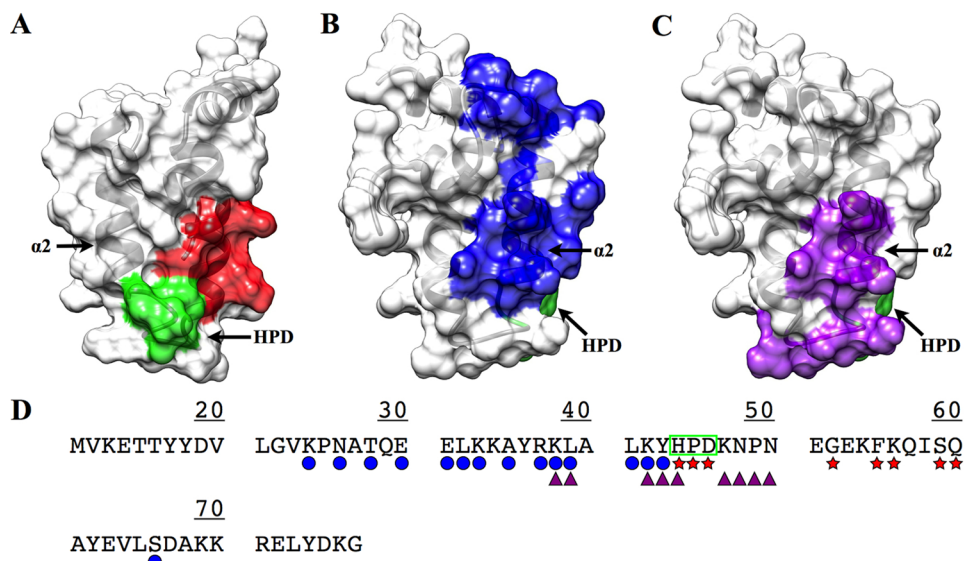


Figure 5. (A) Transparent surface and ribbon representation of DNAJA1-JD highlighting another proposed DnaK binding site based upon the bovine auxilin–bovine Hsp70 complex (PDB entry 2QWN) (colored red and green; conserved HPD motif). (B) Transparent surface and ribbon representation of DNAJA1-JD (rotated $\sim 90^\circ$) with the proposed DnaK binding site based upon NMR titration data (colored blue). (C) Transparent surface and ribbon of DNAJA1-JD (rotated $\sim 90^\circ$) with the proposed inhibition site based upon the TIM14–TIM16 complex (colored purple). (D) Sequence of DNAJA1-JD with the proposed interaction sites indicated: DnaK binding site from titrations (blue circle), DnaK inhibition site (purple triangles), DnaK binding site from the cross-linked auxilin–Hsp70 complex (red stars), and the highly conserved HPD motif (green box). The conserved HPD motif and helix $\alpha 2$, which is potentially an important component of the DnaJ–DnaK interaction site and the TIM16-like inhibitory binding site, are labeled.

the perturbed residues in the *E. coli* DnaJ J-domain occurred along helix $\alpha 2$. An alignment of the *E. coli* DnaJ J-domain and human DNAJA1-JD sequences allowed us to map those same

perturbed residues onto the human DNAJA1-JD sequence and structure (Figure 5B). This process identifies a possible DnaK binding site on DNAJA1-JD that includes helix $\alpha 2$ instead of

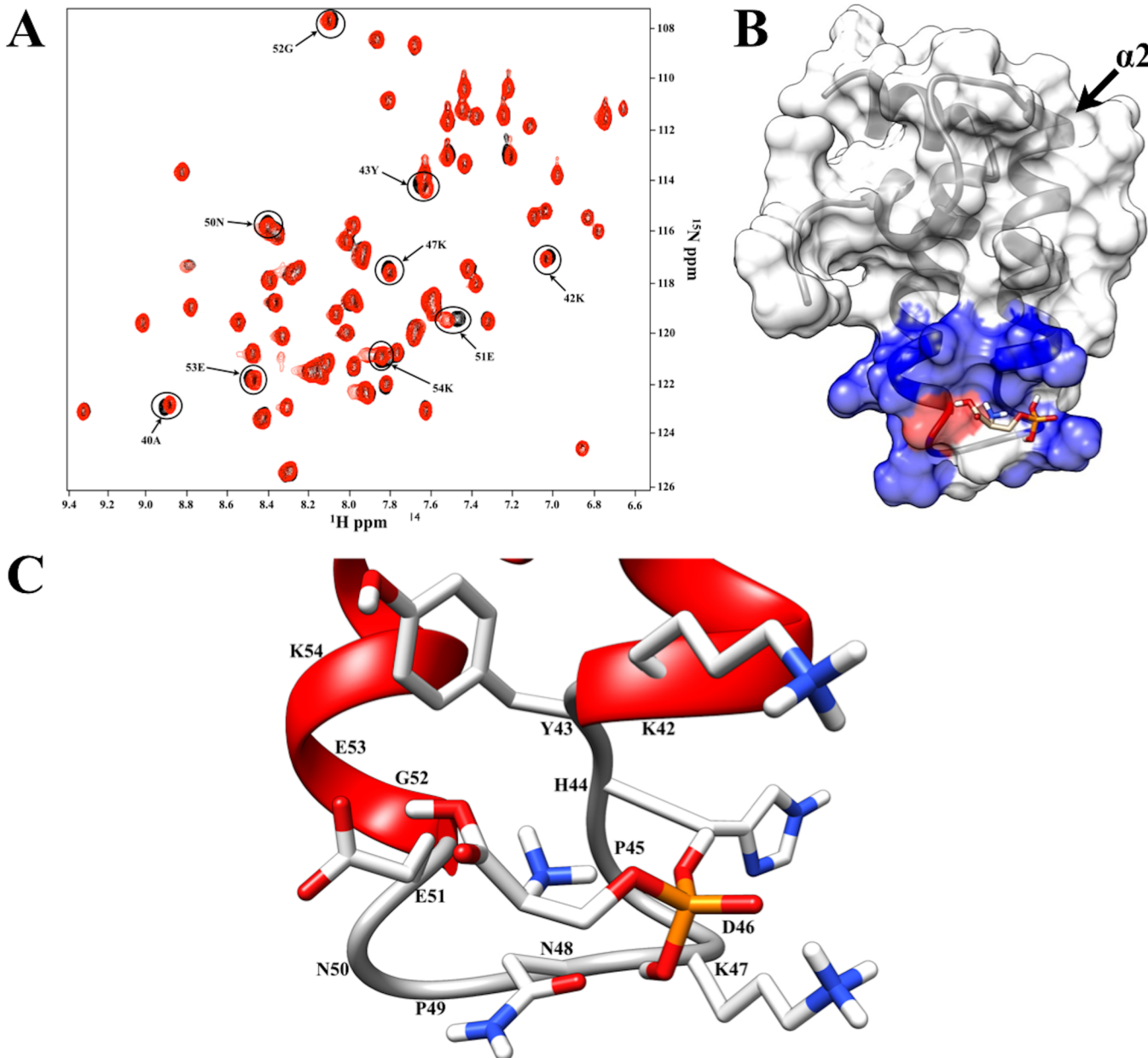


Figure 6. (A) Overlay of 2D ¹H-¹⁵N HSQC spectra of free DNAJA1-JD (black) and DNAJA1-JD with O-phospho-L-serine (red). (B) Transparent surface representation and ribbon diagram of DNAJA1-JD bound with O-phospho-L-serine, with the residues showing a chemical shift perturbation upon binding of O-phospho-L-serine colored blue and the one residue, Glu51, that shows the greatest chemical shift perturbation with every binding ligand colored red. (C) Expanded view of O-phospho-L-serine bound to a ribbon diagram of DNAJA1-JD, where residues with side chains directed toward the ligand are displayed.

the HPD motif predicted from the DnaJ-DnaK X-ray structure. Helix α 2 is an intriguing location for a potential protein binding site because it presents a positively charged surface. Coincidentally, the proposed DnaJ binding site on DnaK has a negatively charged surface, which further supports the identification of helix α 2 as a binding site for DnaK. Additionally, mutations of residues in helix α 2 inhibit the DnaJ-DnaK interaction, further implicating the importance of helix α 2 in binding DnaK.^{90,92}

Another potential protein interaction site on DNAJA1-JD may be linked to the inhibition of DnaJ activity. TIM14 is a related DnaJ protein that is essential for the transport of proteins across the outer membrane of mitochondria by stimulating ATPase activity of mitochondrial Hsp70.⁹³ Any

mutation in the HPD motif of TIM14 effectively inhibits its activity, which indicates the importance of the HPD motif in the function of DnaJ proteins. Importantly, TIM14 activity is inhibited when TIM14 is bound to TIM16. TIM16 is another J-domain-like protein, but it lacks the HPD motif. The location of the TIM14-TIM16 interaction site partially overlaps with the proposed DnaK binding site that includes residues in helix α 2.⁹⁴ A sequence alignment of TIM14 with DNAJA1-JD indicates an essentially identical overlap (Figure 5C), where four of the 15 residues that make up the proposed DnaK binding site are also part of the potential inhibition site (Figure SD). Thus, the binding of a TIM16-like protein to DNAJA1 would sterically interfere with DnaK binding because of this partial overlap in protein binding sites. Again, a TIM16-like

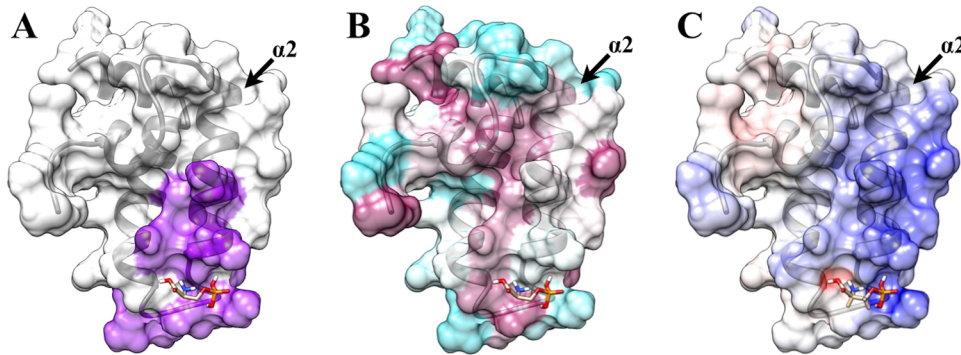


Figure 7. Transparent surface representation and ribbon diagram of DNAJA1-JD bound with O-phospho-L-serine with (A) the proposed inhibition site based on the TIM14–TIM16 interaction (purple), (B) the highly conserved (magenta) and poorly conserved (cyan) residues from Consurf, and (C) the positively charged surface (blue) and negatively charged surface (red) from Delphi. Helix $\alpha 2$, which is potentially an important component of the DnaJ–DnaK interaction site and the TIM16-like inhibitory binding site, is labeled.

protein that binds DNAJA1 has not been identified. Thus, the DnaK binding site is located in the highly conserved HPD motif or helix $\alpha 2$, where a potential inhibitory binding site also involves helix $\alpha 2$. Our ligand binding assay may provide some further insight regarding these potential protein binding sites on DNAJA1-JD.

Identification of a Ligand Binding Site on the DNAJA1 J-Domain.

A high-throughput NMR ligand affinity screen^{66,67} was performed with a function-based compound library.^{64,65} The library contains ~ 460 biologically active compounds consisting of known drugs, protein inhibitors, metabolites, substrates, carbohydrates, fatty acids, hormones, and cofactors. The 1D ^1H line broadening screen identified 27 possible binders. A 2D ^1H – ^{15}N HSQC screen identified only seven compounds that induced chemical shift perturbations (CSPs) upon binding with DNAJA1-JD: O-phospho-L-serine, pyridoxal 5'-phosphate, Bay 11-7082, β -NADPH, ureidosuccinic acid, 2-aminophenol, and D-glucosamine. The seven compounds induced CSPs to the same set of residues, inferring a consistent and unique ligand binding site. Most of the compounds are small, and five of the compounds have a phosphate group or a carboxylic acid group, both of which are likely to be deprotonated at pH 7.0, leaving negatively charged molecules. Importantly, the chemical library contained other compounds with a phosphate or carboxylic acid groups that did not bind DNAJA1-JD, eliminating the likelihood of a nonspecific binding interaction driven only by charge interactions. No other common chemical motif or scaffold is apparent.

Of the seven compounds, O-phospho-L-serine had the greatest number of CSPs (nine) in the 2D ^1H – ^{15}N HSQC screen (Figure 6A). Using the size of O-phospho-L-serine and our CSP-Consensus program, every residue with an observed CSP was determined to be part of a consensus ligand binding site. A DNAJA1-JD–O-phospho-L-serine costructure was then determined on the basis of these CSPs using AutoDock and AutoDockFilter, which identifies the docked pose that best matches the chemical shift perturbation data.⁷³ The costructure selected by AutoDockFilter (Figure 6B) had an AutoDock binding energy of -2.66 kcal/mol, which fits with the average AutoDock binding energy of -2.54 ± 0.40 kcal/mol for all of the docked poses. The costructure has nine residues with any atom within 6 Å of the docked ligand: Ala40, Leu41, Lys42, Tyr43, His44, Lys47, Asn48, Glu51, and Lys54 (Figure 6C). Six of these residues were perturbed in the 2D ^1H – ^{15}N HSQC spectrum (Figure 6A). However, there may be some differences

between the predicted costructure and the actual complex in solution because of local structural rearrangements upon ligand binding. This is likely to occur because AutoDock used a static model of DNAJA1-JD.

On the basis of the protein–ligand costructure, the binding site coincides with the predicted inhibition site based on the TIM14–TIM16 complex (Figure 7A). The O-phospho-L-serine binding site is also consistent with the locations of chemical shift perturbations observed for the other compounds shown to bind in the FAST-NMR assay.^{66,67} Surprisingly, this ligand binding site is not particularly well conserved (Figure 7B). This fact would benefit a drug discovery effort that targets the proposed DNAJA1–TIM16-like complex. While the J-domain is a common fold, the predicted DNAJA1–TIM16-like interaction site appears to be sequence specific, which means drug selectivity can be achieved. Because most of the binding ligands have negatively charged groups, it seems likely that the compounds have an electrostatic interaction with the positively charged region on helix $\alpha 2$ (Figure 7C). Additionally, most of the protein surface, including the binding site, consists of hydrophilic residues, indicating that hydrophobic interactions are unlikely to be energetically favorable. These properties may explain why most of the ligands shown to bind to DNAJA1-JD were small and negatively charged. Nevertheless, these results provide further support for the existence of an inhibitory protein binding site on DNAJA1-JD and the importance of helix $\alpha 2$ to the function of DNAJA1-JD. It is important to note that the ligands shown to bind DNAJA1-JD have a low binding affinity ($K_D > 100$ to $50 \mu\text{M}$), are unlikely to inhibit protein–protein interactions, and have not been shown to inhibit DNAJA1-JD protein binding or to be biologically active in pancreatic cancer cells. Instead, they are simple chemical probes of DNAJA1-JD functional epitopes and provide experimental evidence that supports the identification of the predicted inhibitory protein binding site on DNAJA1-JD.

CONCLUSIONS

Cell-based functional assays showed that the overexpression of DNAJA1 suppresses the stress response capabilities of the oncogenic transcription factor, c-Jun, which is often overexpressed and hyperphosphorylated in cancer cells. c-Jun is part of the JNK signaling pathway, and its phosphorylation state can promote apoptosis or cell proliferation. DnaK has previously been shown to suppress the JNK pathway, which inhibits the hyperphosphorylated, anti-apoptosis state found in pancreatic

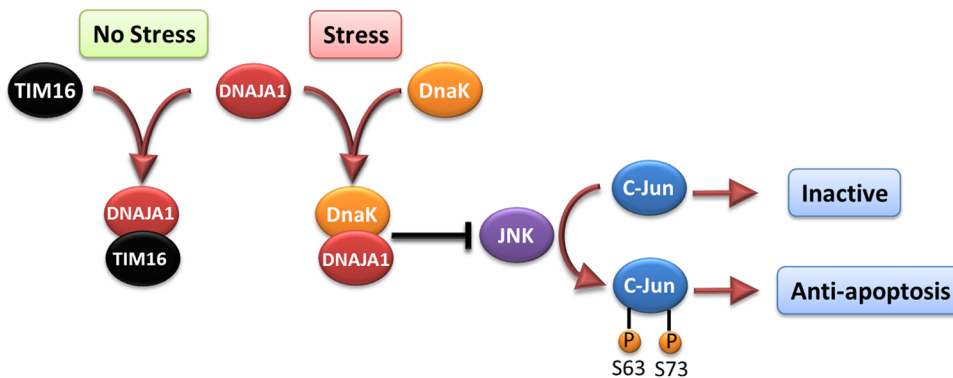


Figure 8. Illustration of the proposed role of DNAJA1 and DnaK on the JNK pathway and c-Jun phosphorylation. The activation of DNAJA1 through the interaction with DnaK appears to suppress the JNK pathway, thus keeping c-Jun in the inactive state. However, inhibiting binding of DNAJA1 to DnaK, as TIM16 does, would allow for the hyperphosphorylation of c-Jun, which has an anti-apoptotic effect.

cancer cells. The DNAJA1-JD NMR structure combined with a bioinformatics analysis and a ligand affinity screen identified a potential DnaK-like binding site and TIM16-like inhibitory binding site. Interestingly, both of these binding sites involve conserved helix $\alpha 2$ and suggest DNAJA1 activity is highly regulated. The NMR ligand affinity screen provided further experimental support for the existence of the TIM16-like inhibitory binding site. On the basis of these observations, we propose a stress response pathway involving DNAJA1 (Figure 8). In the absence of stress, DNAJA1 would be inactivated by forming an inhibitory complex with a TIM16-like protein. In the presence of environmental stress, DNAJA1 would activate a DnaK protein by forming a complex that suppresses the JNK pathway, the hyperphosphorylation of c-Jun, and the anti-apoptosis state. Conversely, the downregulation of DNAJA1 in pancreatic cancer cells likely lowers the activity of DnaK even under conditions of stress, which allows for the hyperphosphorylation of c-Jun and the cell proliferation that is a hallmark of cancer. Thus, the overlapping protein binding interfaces on DNAJA1-JD may be interesting targets for future drug discovery efforts related to pancreatic cancer. Specifically, a drug that disrupts the predicted DNAJA1-TIM16-like inhibitory complex may activate DnaK even in the absence of a stress response. Thus, a constitutively active DnaK may lead to a cascade of events resulting in cell death through the inactivation of JNK and a decrease in the level of c-Jun phosphorylation. Because the overexpression of DNAJA1 reduces pancreatic cancer cell survivability under stress, the J-domain of DNAJA1 itself may be a valuable biological target for treating pancreatic cancer as part of a combination therapy.

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Notes

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ABBREVIATIONS

2D, two-dimensional; 3D, three-dimensional; ATM, auto tune and match; CSPs, chemical shift perturbations; DNAJA1, human protein Dnaj homologue subfamily A member 1; DNAJA1-JD, J-domain of DNAJA1; DSS, 4,4-dimethyl-4-silapentane-1-sulfonic acid; HPD, His-Pro-Asp; HSQC, heteronuclear single-quantum coherence; MTT, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; NESG, Northeast Structural Genomics.

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