

Gene name	NEK1
Uniprot ID	<u>Q96PY6</u>
Region	M1 – K328

Description	NIMA related kinase 1
Synonyms	ALS24, NY-REN-55, SRPS2, SRPS2A, SRTD6
Construct ID	NEK1A-c115
Protein sequence	SMEKYVRLQKIGEGSFGKAILVKSTEDGRQYVIKEINISRMSSKEREESRREVAVLANMKHPNIVQYRESFEENGSLYIVMDYCEGGDLFKRINAQKGVLFQEDQILDWFVQICLALKHVHDRKILHRDIKSQNIFLTKDGTVQLGDFGIARVLNSTVELARTCIGTPYYLSPEICENKPYNNKSDIWALGCVLYELCTLKHAFEAGSMKNLVLKIISGSFPPVSLHYSYDLRSLVSQLFKRNPRDRPSVNSILEKGFIAKRIEKFLSPQLIAEEFCLKTFSKFGSQPIPAKRPASGQNSISVMPAQKITKPAAKYGIPLAYKKYGDKKSSKGGYGLNDIFEAQKIEWHE

Protein mass	with biotin, + 226.1	Protein pl	Extinction (M ⁻¹ cm ⁻¹)
39805.1	40031.1	9.26	38850

Expression host	E. coli BL21(DE3)-R3-lambda-PPase-BirA
NOTE:	To obtain soluble protein from E. coli, NEK1 has to be co-expressed with lambda phosphatase
Expression medium	Terrific broth (TB) with 50 μ g/ml kanamycin, 50 μ g/ml Streptomycin and Chloramphenicol (34 μ g/ml) included in starter cultures; chloramphenicol omitted in final growth.
	Biotin (10 mM stock): dissolve 24mg of d-biotin in 10 ml of warm (microwaved) 10mM bicine buffer (pH 8.3) and sterilize of the solution with a syringe and a 0.2 micron filter.
Transformation and storage	Cells were transformed with the expression plasmid and selected on LB- agar plates containing kanamycin, streptomycin and chloramphenicol. Several colonies were picked from the overnight transformation and used to inoculate cultures in LB supplemented with KAN/Cam/Str which were grown overnight and stored as glycerol (15% v/v) stock at -80°C.
Expression and <i>in vivo</i> biotinylation	10ml starter cultures in LB supplemented with KAN/Cam/Str were grown overnight at 37°C with shaking at 250rpm. Next day, 10 ml was used to inoculate 1L of TB in a 2LUltraYield flask supplemented with Streptomycin 50µg/ml and Kanamycin 50µg/ml. Note: the medium need not contain chloramphenicol at this step; the pRARE2 plasmid is not lost during the growth process and also ensures a more predictable growth. Seal flask with an adhesive air porous film during cell growth. Culture was grown in a 37°C incubator with shaking at 200rpm. Cell growth was monitored until



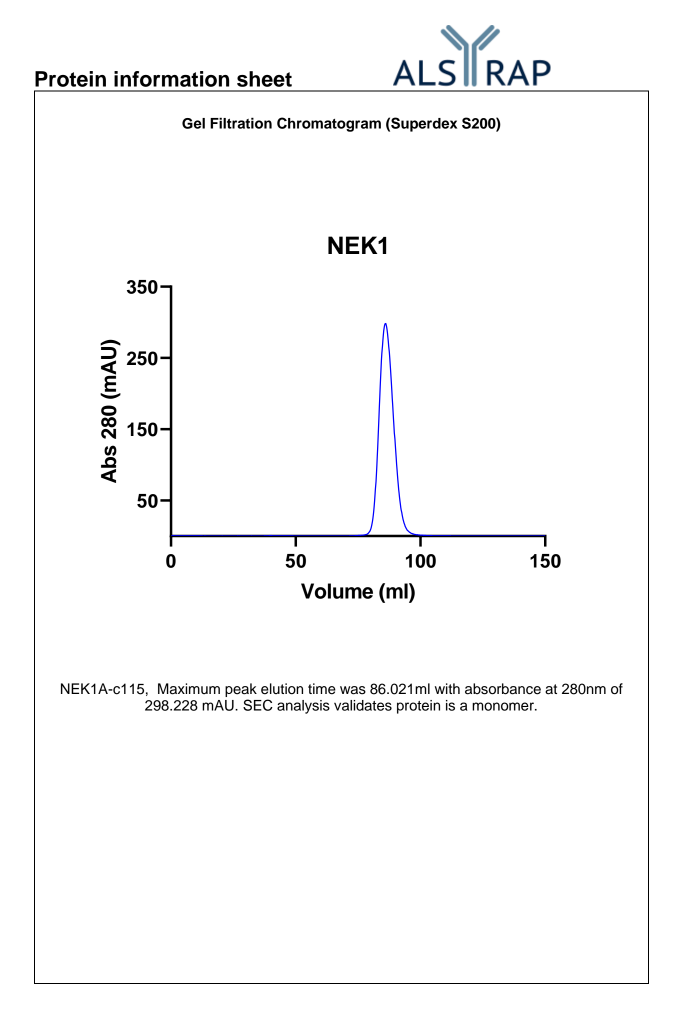
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	mid-log phase was reached and cell density OD_{600nm} was approximately 2.00 – 3.00 in TB. Incubator temperature was lowered to 18°C and culture flask allowed to cool. After 30 - 45 minutes protein expression was induced by the addition of IPTG at 0.1mM final concentration and Biotin solution (10 ml/ 1L culture). Incubation of the culture was continued for 16 hours at 18°C. Next day, the cells were collected by centrifugation (15 min, 5,000 RPM in JLA-8.100 rotors). The supernatants were discarded and the cell pellets transferred into 50ml conical polypropylene tubes. Cell pellets were weighed, frozen and stored at -80°C.
Purification buffers	2x lysis buffer: 100 mM HEPES, pH 7.5, 1.0 M NaCl, 10% v/v glycerol, 1mM TCEP, 10mM imidazole. Protease inhibitors (1:200 Merck cocktail IV) were added before use.
	1x lysis buffer: 50 mM HEPES, pH 7.5, 0.5 M NaCl, 5% v/v glycerol, 1 mM TCEP, 10 mM imidazole
	Wash buffer: 50 mM HEPES, pH 7.5, 0.5 M NaCl, 5% v/v glycerol, 1 mM TCEP, 40 mM imidazole
	Elution buffers: 50 mM Hepes pH 7.5, 500mM NaCl, 5% Glycerol, 1mM TCEP, 100mM, 150mM, 250 mM, 500 mM Imidazole, respectively
	GF buffer: 50 mM HEPES, pH 7.5, 0.5M NaCl, 5% v/v glycerol, 1 mM TCEP
	IEX Low Salt Buffer: 50 mM HEPES, pH 7.5, 50 mM NaCl, 5% v/v glycerol, 1 mM TCEP
	IEX High Salt Buffer: 50 mM HEPES, pH 7.5, 1M NaCl, 5% v/v glycerol, 1 mM TCEP
	PEI 5%, pH 7.5: 5% polyethyleneimine (diluted from 50% stock, Sigma) and titrated to pH 7.5 with HCI.
Purification step 1: IMAC	Cell Lysis: Frozen cell pellets from -80°C were thawed by intermittently placing in a 37°C water bath, then transferred directly to ice. Cells were resuspended in pre-chilled lysis buffer by vortex or pipetting gentle until cells were completely in suspension. 1 volume (v/w) of 2x lysis buffer was added, and the cells were thoroughly suspended. After further addition of 3 volumes (3 ml per 1g of pellet) of 1x lysis buffer the cells were broken by sonication. The lysate was then transferred to a 50ml Beckman polypropylene centrifuge tube. Cell debris and DNA from the lysate was precipitated by adding Polyethylenimine (PEI) to the tube from a 5% stock solution to a final concentration of 0.15% and incubated on ice for 15 – 30 mins for precipitation. Cell lysates were clarified by centrifugation using a JA25.50 rotor at 21,500rpm for 60 min at 4°C. The cleared supernatant was collected and filtered using a 0.45um acrodisc syringe filter and transferred to a fresh 50ml-conical polypropylene tube.
	Ni²⁺-IDA IMAC (column 1): A 25ml Econo-Pac polypropylene column (BioRad) was packed with 5ml of Ni-Sepharose resin and equilibrated in 1x lysis buffer. The cell lysate was loaded onto the column and passed through resin at gravity flow. The unbound flow-through was collected, a wash step of 10 column volumes of wash buffer was included and collected as the wash fraction. Protein was eluted in four intermediate fractions with a step gradient of 5ml imidazole elution buffers; 100mM, 150mM, 250mM



	and 500mM inidenals, respectively, each specient sluted fraction was
	and 500mM imidazole, respectively, each gradient eluted fraction was collected. IMAC fractions were analysed by SDS-PAGE gel for expected molecular weight and purity, the selected eluted fractions were pooled together.
	Mass spectrometry characterisation: Identity and biotinylation efficiency was verified by mass spectrometry, using a Quadrupole-Time of flight (Q-TOF) analyser, Biotin quantitation is measured by: MW of biotinylated protein = MW of sequenced Protein + 226 Da (biotinylation modification).
Step 2: Removal of His ₆ tag and Ni ²⁺ -IDA reverse	Enzymatic removal of histidine tag: Separate cleaved protein from any un-cleaved protein by passing over a Ni ²⁺ -IDA column. Cleaved protein may be recovered in the flow-through fractions. Non-specific contaminating proteins with any un-cleaved protein bound to the column can be eluted with imidazole.
binding	Selected, pooled eluted fractions from Ni ²⁺ -IDA IMAC were concentrated using a 30,000 Amicon-Ultra MWCO centrifugal concentrator to approximately 10mls in volume. Cleavage of the histidine tag was performed by the addition of TEV protease in a 1:20 w/w ratio to the pooled eluted protein fractions. The protein sample plus TEV was transferred to Snakeskin dialysis tubing (3,500 MWCO) and dialysed in 1L Gel Filtration buffer overnight at 4°C on a magnetic stirrer plate for digestion. Next day, the protein was applied to a Ni ²⁺ -IDA reverse-binding column.
	Ni²⁺-IDA Rebinding column (Column 2): To a 25ml Econo-Pac polypropylene column (BioRad), 2ml of packed Ni-Sepharose resin was added, resin was washed in 3 column volumes of distilled deionized water then equilibrated in 5 column volumes of gel filtration buffer. The overnight protein plus TEV sample was loaded and passed through the column by gravity flow. The flow through was collected, to wash any remaining unbound protein from the column a 2x wash step of 2 column volumes of gel filtration buffer was included. Any bound protein was eluted with 2 column volume washes of 40 mM and 500mM imidazole elution buffer. Collected fractions were analysed for verification of cleaved protein and purity by SDS-PAGE gel and mass spectrometry characterisation.
Step 3: SEC: Gel filtration	Size Exclusion Chromatography (SEC) (Column 3): Verified protein from rebinding column 2 was concentrated to required volume for gel filtration (<5 ml), using an Amicon-Ultra 30,000 MWCO centrifugal concentrator. The GF column (Superdex S200 16/60 HiLoad, 120ml on an ÄKTA Express chromatography system) was equilibrated in gel filtration buffer. The concentrated protein sample was loaded onto the GF column for further purification; protein was fractionated on the column in GF buffer at 1ml/min and 1ml fractions were collected at the A280 peaks. Peak fractions were analysed by SDS-PAGE gel, >50% impurities and double bands were observed after gel analysis. The double bands may be due to some protein degradation. Selected GF fractions containing high protein concentration and of a higher purity were collected, pooled and verified using mass spectrometry as one protein species and of the correct mass without any post-translational modifications. Therefore, an additional second GF step was performed attempting to increase purity of the protein. A single peak was observed on the second GF chromatogram, overall



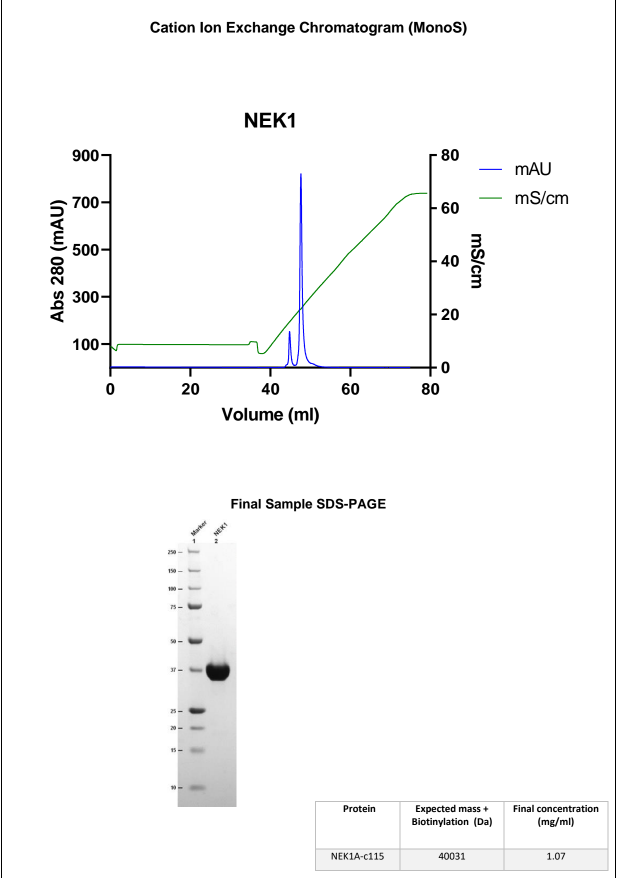
	protein purity was improved but some impurities and degradation remained abound, \geq 30% of total protein sample. Selected peak fractions after a second GF purification containing high protein concentration and purity were collected and pooled and an additional chromatography IEX column was performed for further separation.
Step 4: IEX	Cation Ion Exchange Chromatography (IEX) (Column 4): The IEX column (MonoS 15/50GL, 1ml on an ÄKTA Pure chromatography system) was equilibrated in low salt buffer. The selected peak fractions from the second SEC (GF) were exchanged by dilution into a low salt buffer and loaded onto the column using a super loop The sample load was washed through the column, collecting any unbound protein material. Protein was eluted using a 100% increasing gradient of 50 mM to 1M NaCl at 0.6ml/min over 30CV. Peak fractions were collected and analysed by SDS-PAGE gel. Protein purity was improved after additional IEX step with separation of degradation products and elimination of any bound contaminating non-specific proteins with protein purity increasing to >90%.
Step 5: Protein storage	Protein concentration mg/ml and storage: Protein was concentrated using a 30,000 Amicon-Ultra MWCO centrifugal concentrator and final protein concentration was determined from the absorbance at 280nm, measured using a nanodrop spectrometer (Nanodrop ND-1000, NanoDrop Technologies). Final protein concentration was measured at 1.07mg/ml with a final volume of 2.8ml. Protein was transferred into 200 µl aliquots, flash-frozen in liquid nitrogen and stored at -80°C.



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NEK1A-c115 pNIC-Bio3 40031 40031.6 100% Biotinylated ESI Scan (rt: 0.855 min) Frag=250.0V NEK1 final no tag.d Deconvoluted (Isotope Width=11.9) 40031.63	+ESI Scan (rt: 0.855 min) Frag=250.0V NEK1 final no tag.d Deconvoluted (Isotope Width=11.9)
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