N-Acetyl-D-Glucosamine Kinase Is a Component of Nuclear Speckles and Paraspeckles

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Protein O-GlcNAcylation, dictated by cellular UDP-N-acetylglucosamine (UDP-GlcNAc) levels, plays a crucial role in posttranslational modifications. The enzyme GlcNAC kinase (NAGK, E.C. 2.7.1.59) catalyzes the formation of GlcNAc-6-phosphate, which is a major substrate for the biosynthesis of UDP-GlcNAc. Recent studies have revealed the expression of NAGK in different types of cells especially in neuronal dendrites. Here, by immunocytochemistry (ICC) and immunonucleochemistry (INC) of cultured rat hippocampal neurons, HEK293T and GT1-7 cells, we have showed that NAGK immuno-reactive punctae being present in the nucleoplasm colocalized with small nuclear ribonucleoprotein-associated protein N (snRNPN) and p54NRB, which are speckle and paraspeckle markers, respectively. Furthermore, NAGK IR cluster was also found to be colocalized with GTF2HS (general transcription factor IIH, polypeptide 5) immuno reactive punctae. In addition, relative localization to the ring of nuclear lamin matrix and to GlcNAc, which is highly enriched in nuclear pore complexes, showed that NAGK surrounds the nucleus at the cytoplasmic face of the nuclear outer membrane. By in situ proximity ligation assay (PLA) we confirmed the colocalization of NAGK with snRNPN in the nucleus and in dendrites, while we also verified the interactions of NAGK with p54NRB, and with GTF2HS in the nucleus. These associations between NAGK with speckle, paraspeckle and general transcription factor suggest its regulatory roles in gene expression.

INTRODUCTION

Sugar kinases, stress-70 proteins, and actin belong to a super family defined by a fold consisting of two domains with the topology beta beta alpha beta alpha beta alpha. These enzymes catalyze ATP phosphoryl transfer or hydrolysis coupled to a large conformational change in which the two domains close around the nucleotide (Hurley, 1996). N-acetylglucosamine kinase (GlcNAc kinase or NAGK; E.C. 2.7.1.59), an enzyme of the sugar-kinase/Hsp70/actin super family (Berger et al., 2002), was first identified in 1970 (Datta, 1970). NAGK is a prominent enzyme in amino sugar metabolism where it catalyzes the conversion of GlcNAc to GlcNAc-6-phosphate. This metabolic pathway eventually yields to UDP-GlcNAc, which is utilized in the synthesis of O-/N-glycans and sialic acid. UDP-GlcNAc is also a substrate for O-GlcNAc transferase, which modifies cytosolic and nuclear proteins at serine or threonine residues by adding a single GlcNAc. Furthermore, these O-GlcNAc modifications are associated with regulatory functions ranging from transcription, translation, cell signaling, and stress response to carbohydrate metabolism (Hart et al., 2011; Wells et al., 2003; Zachara et al., 2004).

Murine NAGK is a homodimeric enzyme of 39 kDa subunits (Hinderlich et al., 1998), whereas cDNA sequence of human NAGK encodes a protein of predicted molecular mass 37.4 kDa (Hinderlich et al., 2000). Crystal structures of homodimeric human NAGK revealed that the N-terminal small and C-terminal large domains form a V-shaped structure which acts as an active center for binding GlcNAc and ATP (Weihofen et al., 2006). Recently, our laboratory reported a non-canonical function of NAGK in neuronal dendritogenesis (Lee et al., 2014a). It was found that NAGK is highly expressed in neuronal dendrites, and NAGK overexpression causes significant increase in the complexity of dendritic arborization, such as, increases in the number of primary dendrites and dendritic branches. In contrast, knockdown of NAGK by short hairpin RNA caused significant dendritic degeneration. In a following article, our laboratory further showed that this non-canonical function is not dependent on the kinase activities of NAGK (Lee et al., 2014b). During these studies, predominant NAGK clusters were noticed in the nuclei of all types of cells observed, which included neurons, astrocytes, human embryonic kidney cell-line HEK293T and mouse hypothalamic GT1-7 cell line. Multiple domains exist in nucleoplasm, and the elucidation of the subnuclear domain where NAGK locates would provide insights regarding its nuclear function. Toward this end, we employed immunocytochemistry (ICC), immunonucleochemistry (INC; Moon et al., 2010), and a proximity ligation assay (PLA), to determine the nuclear location of NAGK.

MATERIALS AND METHODS

Antibodies

The following antibodies were used at the indicated dilutions: chicken polyclonal N-acetylglucosamine kinase (cNAGK) (1:1,000;
GenWay Biotech, Inc., USA); O-linked N-acetylgalactosamine (GlcNAc_RL2) (1:1000; Affinity BioReagents Inc., USA); MAb actin (1:500; Sigma), rabbit polyclonal N-acetylgalactosamine kinase antibody (1:500; GeneTex, Inc., USA); mouse monoclonal GlcNAc kinase (1:50; Santa Cruz Biotechnology); mouse monoclonal (MAb) tubulin α-subunit (α-Tubulin) (1:2000; Developmental Studies Hybridoma Bank, University of Iowa, USA); mouse MAb promyelocytic leukemia (PML) (1:10; PG-M3; sc-9666; Santa Cruz Biotechnology); rabbit polyclonal Nucleolin (1:1000; Sigma); rabbit polyclonal snRNPN (1:1000; Sigma); mouse MAb p54NRB (1:1000; BD Biosciences, Inc., USA); rabbit polyclonal lamin B (1:1000; Young In Frontier Inc., Korea); rabbit Anti-p8/TTD-A (GTF2H5) (1:100; Sigma). The fluorescently labeled secondary antibodies Alexa Fluor 488, 568 (Invitrogen, OR) were used to detect the primary antibodies.

Primary culture of rat hippocampal neurons

Hippocampi from embryonic day 19 (E19) Sprague-Dawley rat pups were dissected, dissociated by trypsin treatment and mechanical trituration, and plated onto 12-mm diameter poly-DL-lysine/laminin coated glass coverslips at a density of ~150 neurons/mm², as previously described (Brewer et al., 1993). Cells were initially plated in MACS® Neuro Medium (Miltenyi Biotec Inc., USA) supplemented with MACS NeuroBrewTM-21 (Miltenyi Biotec Inc.), 45.95 μM glutamate, 500 μM glutamine, 25 μM 2-mercaptoethanol, and 1% penicilllin streptomycin, and fed at 4-day intervals after plating with the same media, but without glutamate and 2-mercaptoethanol. GT1-7 and HEK293T cell lines. Image contrast and brightness were optimized using Adobe Systems Photoshop 7.0 software.

RESULTS

Nuclear expression of NAGK in cultured rat hippocampal neurons

Initially, we applied ICC to investigate the expression of NAGK in cultured rat hippocampal neurons. As revealed by double-labeling with anti-NAGK and anti-tubulin (Fig. 1A-a) or anti-actin (Fig. 1A-b) antibodies, NAGK-immunoreactive (IR) signals were distributed in the somatodendritic domain and nucleus. In dendrites, the distribution of NAGK was very similar to that of tubulin (Fig. 1A-a). Strong IR signals for both NAGK and tubulin were associated with dendritic shafts (asterisks) while the

![Fig. 1. Nuclear expression of NAGK in hippocampal neurons. Cultured rat hippocampal neurons (DIV 21) were double-labeled with indicated antibodies. (A) ICC. Double-labeling with anti-NAGK/tubulin (a) or with anti-NAGK/actin (b) revealed the expression profile of NAGK was similar to that of tubulin in dendroplasm (asterisks). NAGK clusters in the central area of soma (arrows) and a circular arrangement (arrowheads) were marked. (B) INC. Double-staining ‘naked’ nuclei with anti-NAGK/tubulin antibodies revealed patch-like NAGK-IR signals in nucleoplasm (arrows) with a circular distribution (arrowheads). The boxed area is enlarged in the inset on the right. Scale bar; 10 μm.](http://molcells.org)
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NAGK and actin-IR signals were not well colocalized; actin-IR signals were distributed at the periphery while NAGK-IR signals were concentrated in the shafts of dendrites (Fig. 1A-b). This feature of NAGK distribution in dendrites implies that NAGK plays a role in dendritic shafts in association with microtubules. In addition to dendritic shafts, NAGK-IR signals also formed clusters in the central area of soma (arrows). These NAGK clusters were roughly round with diverse sizes, the larger ones being ~1-2 μm in diameter. Occasionally in ICC images, small NAGK clusters formed a circle in soma with a diameter close to that of the nucleus (arrowhead in Fig. 1A-b). These features indicate that NAGK is present in the nucleoplasm and in the nuclear envelope.

INC visualized the localization of NAGK in the nuclear envelope
The circular distribution of NAGK in soma was not always evident by conventional ICC, probably due to surrounding constituents in soma. Therefore, we applied the INC, which involves the immunostaining of 'naked' nuclei after the removal of cytoplasmic components (Moon et al., 2010), to clearly observe circular NAGK IR signals in the soma. Cultured rat hippocampal neurons were subjected to double labeling with antibodies against NAGK and tubulin. The INC process removed most of the cytoplasmic components, including NAGK, as was observed in a fluorescence image (Fig. 1B). Only residual amounts of dendritic tubulin-IR signals, which resisted dissolution due to tight microtubule bundling in neuronal dendrites, were left (Fig. 1B, asterisk). In this image nuclei lit up like moons and the circular distribution of NAGK IR signals was clearly observed around nuclei (arrowheads; Fig. 1B, inset) and large NAGK clusters were observed in nucleoplasm (arrows). These results demonstrate the presence of NAGK in nucleoplasm and its circular distribution along the nuclear envelope.

NAGK localized at the outer membrane of the nuclear envelope
The nuclear envelope is a double unit membrane separated by intermembrane lumen. In most cells, on the inner surface of the nuclear envelope there is a thin fibrous lamina of variable thickness (Fawcett, 1966). This lamina consists of polypeptides called lamin. In thin sections, these form a meshwork of extremely fine filaments that runs parallel to the inner surface of the envelope. The present study indicates that NAGK forms a circle around nuclear envelope. To pinpoint the locations of these NAGK clusters in relation to the nuclear envelope, we performed INC using markers for the inner and outer surfaces of the envelope. For an inner surface marker we used lamin, which forms a thin matrix along the nucleoplasmic face of inner nuclear membrane. For an outer nuclear membrane marker we used GlcNAc, because the unstructured phenylalanine-glycine (FG) repeat regions of the nuclear pore complex are heavily O-GlcNAc-modified, and extend on both the nuclear and cytoplasmic sides of the pore (Li and Kohler, 2014). Therefore, we first investigated whether GlcNAc could serve as a nuclear envelope marker. For this purpose, we double-labeled HEK293T cells with antibodies against GlcNAc and lamin. GlcNAc-IR signals formed clusters (red arrowhead) outside the lamin circles (Fig. 2A, green arrowhead) indicating that these GlcNAc-IR signals were from the nuclear envelope. Having verified that GlcNAc-IR signals represent the nuclear envelope, we double-labeled HEK293T cells and cultured rat hippocampal neurons with anti-NAGK/-lamin (Fig. 2B) or anti-NAGK/-GlcNAc (Fig. 2C) antibodies. The merge images of NAGK- and lamin-IR signals revealed that NAGK clusters lie beyond lamin rings (Fig. 2B, red arrowheads). Notably, in many cases NAGK-IR signals were completely separated from lamin rings (Fig. 2B-b, arrowheads) indicating that the NAGK-IR ring lies well beyond the lamin-IR ring. The merged double-labeled image of NAGK and GlcNAc revealed NAGK rings at the outer margin of GlcNAc-IR signals, which represent the cytoplasmic face of the outer nuclear membrane. Scale bar; 10 μm.

NAGK localized to nuclear speckles and paraspeckles but not to PML body or nucleolus
The nucleoplasm is compartmentalized into highly dynamic sub
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NAGK localization at nuclear subcompartments. Double-labeling of rat hippocampal neurons (DIV 21) with anti-NAGK and various markers for subnuclear domains shows that NAGK colocalized well with snRNPN (a speckle marker; A) and with some select paraspeckles (B-a). Colocalization of NAGK with the paraspeckle marker p54NRB was partial (yellow arrowheads) or absent (arrows) in HEK293T cells (B-b). (C) Immunonucleochemistry of GT1-7 cell showed that General transcription factor 2H5 or TTDA overlay on NAGK-IR clusters, (inset area showing the overlaid portion). NAGK did not colocalize with PML protein (D) or nucleolin (E; a marker for nucleoli). Scale bar; 10 μm.

NAGK colocalized with snRNPN also in neuronal dendrites
Considerable evidence supports that pre-mRNA splicing takes place in neuronal dendrites. For example, some auxiliary constituents of spliceosome, like SMN (Jurica and Moore, 2003) and SAM68 (Grange et al., 2004), were found throughout neuronal cytoplasm and to extend to dendrites. Furthermore, the pre-mRNA splicing factors found in dendroplasm were reported to retain the potential to promote pre-mRNA splicing in isolated dendrites (Glanzer et al., 2005). The present and our previous study show that NAGK is highly expressed in neuronal dendrites (Lee et al., 2014a, b). Therefore, we performed ICC to investigate the possibility of NAGK-snRNPN colocalization in dendroplasm. Accordingly, we double-labeled cultured hippocampal neurons (DIV 6) with anti-NAGK and anti-snRNPN antibodies, and resulting fluorescence images showed that both NAGK- and snRNPN-IR punctae were interspersed throughout dendrites (Fig. 4). Merged images revealed the presence of some colocalization of these two proteins (Fig. 4C, arrowheads in inset), suggesting a direct interaction between them.

NAGK interacted with snRNPN, p54NRB and GTF2H5
Our initial trials to find the proteins interacting with the small domain of NAGK by yeast two-hybrid selections screened out two yeast clones (clone #3 and 5) coding for snRNPN; a paralog of the snRNPB (B) and paraspeckle (p54NRB). The colocalization of NAGK with snRNPN in neurons (Fig. 3A) and with snRNPN in HEK293T cells (not shown) was almost perfect. The colocalization profile of NAGK with a paraspeckle (p54NRB) marker was varied depending on cell types (Fig. 3B). In neurons, NAGK and p54NRB immunopunctae partially overlapped (Fig. 3B-a), whereas in HEK293T cells, NAGK clusters colocalized partially, as in neurons, with p54NRB clusters in some cells (yellow arrowheads). On the other hand, the two proteins were completely segregated into different domains in other HEK293T cells (Fig. 3B-b, arrows). INC of GT1-7 cells revealed GTF2H5-IR signals overlaid on NAGK clusters (Fig. 3C). NAGK clusters did not colocalize with markers for PML body (Fig. 3D) or nucleolus (nucleolin) (Fig. 3E).
were added, showed no specific PLA signal both in neurons and GT1-7 nucleus (data not shown). These results indicated that NAGK directly interacts with snRNPN, p54NRB, and GTF2H5.

**DISCUSSION**

In this study, we describe the characteristic distribution patterns of NAGK in different sub nuclear compartments. ICC of cultured hippocampal neurons showed somatodendritic expression of NAGK, and its nuclear expression profile was clearly depicted by INC where an irregular cluster-like expression pattern in nucleoplasm and a circular arrangement along the outer side of the nuclear envelope was found. Double-labeling of ‘naked’ nuclei in INC showed that NAGK was colocalized with snRNPN (a speckle marker protein), p54NRB (a paraspeckle marker protein), and TTDA (GTF2H5). We also found these protein interactions (snRNPN, GTF2H5) by PLA and in yeast two hybrid screening.

Subcompartments of nucleoplasm are involved in the splicing, translation, and transcriptional regulation of genes (Spector, 2001; Zimber et al., 2004). Of these, speckle (also known as interchromatin granule cluster) is a key nuclear subcompartment where splicing and transcription take place simultaneously. Speckle is considered the hub for many pre-mRNA splicing factors, including snRNPs and serine/arginine-rich (SR) proteins (Fu, 1995). Of the snRNPs, snRNAs, such as, U1, U2, U4, U5 and U6 (7), form complexes with B, B', D, D', E, F and G snRN associated proteins. snRNPN is a paralog of the snRNPB'/B gene, which is alternatively spliced to produce SmB or SmB' spliceosomal core proteins (Gray et al., 1999). snRNPN is expressed in a tissue-specific manner, and is most abundant in rat brain, less so in heart and undetectable in other tissues examined (McAllister et al., 1988). Therefore, it is interpreted that an ancestral duplication of the snRNPB'/B gene gave rise to the snRNPN paralog which evolved new functions in specific tissues. However, our PLA and INC results showed that NAGK interacts with SmB'/B in HEK293T (data not shown) and snRNPN in neurons, which suggests that NAGK has the same function in neuronal and non-neuronal cells.

In addition to pre-mRNA splicing, mounting evidence suggests that speckle also modulates transcription processes. For examples, a subpopulation of the largest subunit of RNA polymerase II molecules is distributed in speckle domains in transcriptionally active nuclei (Bregman et al., 1995). Significant levels of RNA Polymerase II-mediated transcription sites are associated with nuclear speckles (Wei et al., 1999). Depletion of a prototypical SR protein SC35 induced Pol II accumulation within the gene body and attenuated elongation, suggesting speckle proteins play critical roles in transcriptional elongation (Lin et al., 2008). About 80% of pre-mRNA splicing occurs co-transcriptionally, and active spliceosomes localize in situ to regions of decompacted chromatin at the periphery of or within nuclear speckles, whereas the release of post-transcriptionally spliced mRNA from speckles is coupled to the nuclear mRNA export pathway (Girard et al., 2012).

The present study also shows snRNPN/NAGK interaction in neuronal dendrites, indicating involvement of NAGK in local splicing process at dendrites. Splicing in dendrites has been extensively researched since the discovery of protein synthetic machinery including ribosomes, membranous constituents of endoplasmic reticulum, and Golgi apparatus in dendroplasm. Nova proteins (neuron-specific RNA binding proteins) regulate alternative splicing in neurons (Jensen et al., 2000), and immunofluorescence and EM studies of spinal cord motor neurons demonstrated that Nova localizes within synaptic contacts in

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**Fig. 5.** Interaction of NAGK with nuclear proteins. (A) Yeast two hybrid assays using the NAGK small domain as bait identified snRNPN and GTF2H5. The NAGK-interacting domains of snRNPN (a) and GTF2H5 (b) of two different clones for each protein are shown. The positions of like-Sm (LSM) and TTDA subunit of TFIIH basal transcription factor complex (Tbf5) protein domains of each protein were marked on the bars with amino acid numbers. (B) PLA showing interactions of NAGK with nuclear proteins. PLA was performed in rat hippocampal neurons (DIV 11). A low power image shows distribution of NAGK/snRNPN PLA dots (arrowheads) in soma (inset) and dendrites (a). Immunonucleochemistry followed by PLA shows the interaction between NAGK and snRNPN (arrows) in the ‘naked’ neuronal nucleoplasm (b). Scale bars: 5 and 20 μm in a and b, respectively. PLA conducted in HEK293T cells showed interaction between NAGK and paraspeckle protein (p54NRB) in nucleus (c, arrowheads). Scale bar, 5 μm. PLA done in hypothalamic GT1-7 cells showed NAGK/TTDA (GTF2H5) PLA dots in nucleoplasm (arrowheads). Scale bar, 10 μm.
dendrites (Racca et al., 2010), which demonstrates the presence and functionality of pre-mRNA-splicing components in dendrites. For example, splicing factors, SF2/ASF, USAF65, and Sm or SR proteins are located in proximal and distal portions of dendrites and their branch points. Furthermore, reporter pre-mRNAs can be translated in dendritic cytoplasm after dendritic splicing and when isolated dendrites are transfected with chicken delta-crystallin pre-mRNA or luciferase reporter pre-mRNA, splicing junctions clustered at or near expected splice sites (Glanzer et al., 2005). These observations suggest that pre-mRNA-splicing factors in dendroplasm retain the potential to promote pre-mRNA splicing. It is now widely accepted that translation of mRNAs localized to dendrites is regulated in response to neural activity and is required for many forms of long term synaptic plasticity. Thus, it appears NAGK may play a role in the splicing of the premRNAs involved in local dendritic translation. Further research on NAGK is needed to elucidate its role in neuronal dendrites.

In the present study, we found that many NAGK clusters overlapped those of p54nrb, a paraspeckle marker, in neurons and some HEK293T cells. However, in some HEK293T cells, NAGK formed a few large clusters that were completely segregated from p54nrb. Paraspeckles are nuclear bodies that are usually visualized in cultured cell lines as 2–20 foci during interphase and during all of mitosis except for telophase (Naganuma and Hirose, 2013). At telophase, when the two daughter nuclei are formed, there is no RNA polymerase (Pol) II transcription. When Pol II transcription is inhibited, paraspeckle protein 1 (PS1P) leaves paraspeckles and accumulates at the nucleolus, concentrating in crescent-shaped structures termed “perinucleolar caps” (Fox et al., 2002). p54nrb forms a heterodimer with PS1P and is targeted to paraspeckles in a RNA-dependent manner whereas its accumulation at perinucleolar caps does not require RNA (Fox et al., 2005; Shav-Tal et al., 2005). We did not study whether NAGK is present in the perinucleolar cap. However, our study showed that NAGK is not present in the nucleolus. Therefore, in cultured HEK293T cells, some cells in Pol II inhibition may exhibit segregation in NAGK and p54nrb signals. Paraspeckles are involved in diverse gene expression associated regulatory events such as regulation of the transcription and pre-mRNA splicing events associated with important physiological events and diseases (Lagier-Tourenne and Cleveland, 2009; Shav-Tal and Zipori, 2002). NAGK may participate in these processes.

Interaction of NAGK with GF2F2H (TTDA) was confirmed by the yeast two-hybrid assay, INC and PLA. These findings support that NAGK might have a role in gene transcription. GF2F2H, a multi subunit complex, is essential for Pol II transcription and Nucleotide Excision Repair (NER). GF2F2H is a small (8 kDa) protein subunit, of which mutation causes trichothiodystrophy (TTD-A) disease. Under normal condition TTDA maintained equilibrium of free and GF2F2H bound states, but shifts into high GF2F2H bound condition in presence of DNA lesion by UV-irradiation (Giglia-Mari et al., 2006; Nonnekens et al., 2013). Therefore, NAGK may regulate NER by binding with GF2F2H. In summary, we show that NAGK interacts with speckle (snRNPs), paraspeckle (p54nrb), GF2F2H in the nucleolus, and also present in outer nuclear membrane. Altogether, our present study implies a diverse but significant non-canonical function of NAGK in the nucleolus.

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