WNK2 (WNK lysine deficient protein kinase 2)

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Identity

Other names: KAIA302, KIAA1760, NY-CO-43, P/OKcl.13, PRKWNK2, SDCCAG43
HGNC (Hugo): WNK2

Location: 9q22.31

Local order: The WNK2 gene is covered by BAC clones RP11-370F5, RP11-480F4 and RP11-165J3 and is flanked by the NINJ1 (telomeric) and C9orf129 (centromeric) genes.

DNA/RNA

Description

The human WNK2 gene is composed of 31 exons spanning 136 Kbp on chromosome 9q22.31. The promoter region contains a 700 bp CpG island between 1103 bp and 396 bp upstream of the ATG translation start codon. A second CpG island spans exon 1 from 135 bp upstream of the ATG translation start codon until 638 bp downstream of the ATG and close to the end of exon 1.

Transcription

Two major alternative transcripts exist depending on the terminal exon chosen.

Protein

Description

Amino acids: 2297 or 2254. Molecular Weight: 243000 Daltons. The WNK2 protein encodes a cytoplasmic serine-threonine kinase that lacks a lysine in subdomain II required for ATP-binding in most protein kinases and instead uses an alternative lysine in subdomain I. WNK kinase form a separate family branch, most closely related to kinases MEKK, Raf and PAK.

One variant uses exons 1-30, has a coding sequence of 6894 bp and yields WNK2(1-2297) (related to clone KIAA1760, Acc. Nb. AB051547). The other variant skips exon 30 and includes exon 31, has a coding sequence of 6765 bp and yields WNK2(1-2254) (related to clone Kaia302; Acc. Nb. AK000694). Both terminal exons 30 and 31 carry their own 3’-untranslated regions and polyadenylation signals. In addition, there is evidence for alternative splicing in other exons and in a tissue-specific manner.

Pseudogene

None known.

Human WNK2 gene structure. The gene spans 136 Kbp, contains 31 exons and localizes to chromosome 9q22.31. Exons (vertical boxes) and separating introns are shown in proportion to their sizes; however, intron scale differs from exon scale.
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Expression

WNK2 is preferentially expressed in heart, skeletal muscle and brain but also in small intestine, colon and liver. Loss of expression was reported in a large percentage of human gliomas (Hong et al., 2007) and grade II and III meningiomas (Jun et al., 2009) due to extensive methylation in the CpG island at the 5’ end of the WNK2 gene. In contrast, promoter methylation was rare in other tumor types. This finding makes WNK2 a candidate tumor suppressor gene in brain tumors.

Localisation

The subcellular localization of GFP-tagged WNK2 in HeLa cells was predominantly cytoplasmic. Part of the endogenous WNK2 pool in HT29 colorectal cells localized to the plasma membrane and overexpression of a WNK2(1922-2156) that contains the coiled-coil domain was targeted to the plasma membrane.

Function

Human WNK2 modulates the activation level of ERK1 and ERK2. Experimental depletion of WNK2 or overexpression of a kinase-dead WNK2K207M mutant led to increased phospho-ERK1/2 levels when a basal ERK stimulation was present but not, for example, in serum-free culture conditions (Moniz et al., 2007). This increase in ERK1/2 activation promoted cell cycle progression through G1/S and sensitized cells to respond to lower concentrations of EGF. From these data one might predict that loss of WNK2 expression will promote cell cycle progression in tumor cells.

Interestingly, WNK2 expression is silenced in a significant percentage of human gliomas (Hong et al., 2007) suggesting that this pathway may be used in some tumor types to promote cell proliferation. The molecular mechanism through which a reduction in WNK2 expression can increase ERK1/2 activation involves phosphorylation of MEK1 at serine 298, a modification that increases MEK1 affinity towards ERK1/2. Apparently, WNK2 affects PAK1 activation via Rac1 and PAK1 is the kinase responsible for MEK1 S298 phosphorylation (Moniz et al., 2008).

Homology

The catalytic domain of WNK2 is 90% identical to WNK1, 91% identical to WNK3 and 81% identical to WNK4. The remaining sequence of WNK2 has little homology to other WNK members except for three small WNK homology regions (Holden et al., 2004; Moniz et al., 2007). These include an acidic motif (residues 586-597) to which hereditary mutations in WNK4 cluster (Wilson et al., 2001), residues 1186-1261 without any recognizable motif, and residues 1918-1988 including a coiled-coil domain.

Mutations

Note

At present it is unclear whether the observed somatic mutations have a functional impact on the WNK2 protein or confer any selective advantage to tumors cells.
<table>
<thead>
<tr>
<th>Tissue</th>
<th>Histology/Type</th>
<th>cDNA</th>
<th>Protein</th>
<th>Mutation</th>
<th>Ref</th>
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<tbody>
<tr>
<td>Colorectal</td>
<td>adenocarcinoma</td>
<td>c.1964delC</td>
<td>p.P655fs*2</td>
<td>Frameshift deletion</td>
<td>Greenman et al., 2007</td>
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<td>Brain</td>
<td>glioblastoma</td>
<td>c.3799G&gt;A</td>
<td>p.A1267T</td>
<td>Missense</td>
<td>Parsons et al., 2008</td>
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<tr>
<td>Stomach</td>
<td>adenocarcinoma</td>
<td>c.4269delC</td>
<td>p.S1424fs*5</td>
<td>Frameshift deletion</td>
<td>Greenman et al., 2007</td>
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<tr>
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<td>neuroendocrine carcinoma</td>
<td>c.5009G&gt;A</td>
<td>p.G1670E</td>
<td>Missense</td>
<td>Greenman et al., 2007; Davies et al., 2005</td>
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<tr>
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<td>c.6089G&gt;T</td>
<td>p.S2030I</td>
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<tr>
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<td>p.T2267fs*31</td>
<td>Frameshift deletion</td>
<td>Greenman et al., 2007</td>
</tr>
</tbody>
</table>

**Germinal**
No germinal mutations described.

**Somatic**
Somatic mutations in WNK2 have been found in the course of large scale tumor genome sequencing efforts (see Table above). Heterozygous somatic mutations in the WNK2 gene identified by large-scale tumor sequencing.

**Implicated in**
**Brain tumors**

**Note**
Promoter methylation leads to loss of expression.

**Disease**
Glioma and meningioma.

**Prognosis**
Unknown.

**Colon cancer**

**Note**
WK2 clone was isolated as a serologically defined colon cancer antigen 43; WNK2 is expressed in colon.

**To be noted**
Possible role in invasion due to the effect of WNK2 on Rho-GTPases. WNK2 controls (through a yet unknown mechanism) the activation of RhoA, which in turn determines the activation of Rac1 in a reciprocal manner. Experimental depletion of WNK2 leads to reduced RhoA and increased Rac1 activation.

**References**


*This article should be referenced as such:* Jordan P. WNK2 (WNK lysine deficient protein kinase 2). Atlas Genet Cytogenet Oncol Haematol. 2011; 15(1):77-79.