High-Resolution Array CGH Profiling Identifies Na/K Transporting ATPase Interacting 2 (NKAIN2) as a Predisposing Candidate Gene in Neuroblastoma

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Abstract

Neuroblastoma (NB), the most common extracranial solid tumor of childhood, usually occurs sporadically but also its familial occurrence is known in 1-2% of NB patients. Germline mutations in the ALK and PHOX2B genes have been found in a subset of familial NBs. However, because some individuals harbouring mutations in these genes do not develop this tumor, additional genetic alterations appear to be required for NB pathogenesis. Herein, we studied an Italian family with three NB patients, two siblings and a first cousin, carrying an ALK germline-activating mutation R1192P, that was inherited from their unaffected mothers and with no mutations in the PHOX2B gene. A comparison between somatic and germline DNA copy number changes in the affected siblings by a high resolution array-based Comparative Genomic Hybridization (CGH) analysis revealed a germline gain at NKAIN2 (Na/K transporting ATPase interacting 2) locus in one of the sibling, that was inherited from the parent who does not carry the ALK mutation. Surprisingly, NKAIN2 was expressed at high levels also in the affected sibling that lacks the genomic gain at this locus, clearly suggesting the existence of other regulatory mechanisms. High levels of NKAIN2 were detected in the MYCN-amplified NB cell lines and in the most aggressive NB lesions as well as in the peripheral blood of a large cohort of NB patients. Consistent with a role of NKAIN2 in NB development, NKAIN2 was down-regulated during all-trans retinoic acid differentiation in two NB cell lines. Taken together, these data indicate a potential role of NKAIN2 gene in NB growth and differentiation.

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Introduction

Neuroblastoma (NB), the most common extracranial solid tumor of childhood, originates from sympathetic neuronal progenitor cells and accounts for more than 15% of all pediatric cancer deaths [1,2]. The heterogenous clinical behaviour, ranging from localized mass that undergo spontaneous regression, to widespread NB with a survival rate of 50% despite intensive multimodal cytotoxic therapy, is attributable to both biological and genetic characteristics of the tumor [3].

Approximately 1–2% of NB cases have a NB-positive familial history with patients either distributed along three generations or clustered in the most recent one [1]. Mutations in paired-like homeobox 2B (PHOX2B) and the tyrosine kinase receptor anaplastic lymphoma kinase (ALK), two genes predominantly expressed in the developing peripheral nervous system, have been identified as predisposing events for NB. Germline missense mutations in PHOX2B were originally observed in a familial case of NB and in a patient with NB associated with congenital central hypoventilation syndrome (CCHS) and/or Hirschsprung disease [4]. Subsequent studies confirmed the association of PHOX2B mutations with NB predisposition [5–7], and showed that PHOX2B mutations could also be observed as de novo mutations in apparently sporadic tumors [9,9]. More recently, ALK has been demonstrated to be constitutively activated by gene mutations in the tyrosine kinase domain and/or amplification in sporadic [10–13], as well as in familial cases of NB [14,15]. The range of somatic and germline ALK mutations are different, being some mutations detected either in familial or in sporadic forms of NB.

Somatically acquired genomic aberrations are of fundamental importance for predicting NB patients’ outcome. Tumors with amplification and overexpression of the MYCN proto-oncogene or allelic loss of chromosome arms 1p and 11q or both, typically are metastatic at diagnosis and therapy-resistant [1]. Conversely, tumors showing hyper- or hypoploidy due to whole-chromosomal gains or losses, respectively, are more easily cured and frequently undergo spontaneous regression [1].
Although germline mutations of \(\text{ALK}\) and \(\text{PHOX2B}\) account for the majority of familial NB, additional genetic factors responsible for NB tumor progression may still be discovered. Consistently, a recent paper demonstrates that \(\text{ALK}\) deleterious mutations are rare events in NB patients with a high probability of predisposition [16], strongly suggesting that other predisposing genes remain to be identified.

Recently, the application of whole genome high-resolution array comparative genomic hybridization (array CGH) techniques has allowed the identification of small copy number changes in cancer genomes [17]. Taking advantage of this technology, we screened NB-affected and unaffected individuals of an Italian family to systematically identify germline and tumor-specific copy number alterations relevant to NB. We identified 3 new genomic alterations in chromosomal regions devoid of any copy number variations (CNV). One of these genes, \(\text{NKAIN2}\), was detected upregulated in a large cohort of NB patients (both tumor and blood specimens) and NB cell lines with aggressive phenotype, suggesting its putative role in NB pathogenesis.

Results

Neuroblastoma family case

This family includes three NB patients, two siblings and a first cousin (Figure 1). The two siblings were affected by stage 3 and stage 4 NB according to INSS [18], III-2 and III-3, respectively, Figure 1). The sibling III-2 is the eldest daughter of non-consanguineous parents. She was diagnosed at age of 16 months with an abdominal paravertebral \(\text{MYCN}\)-non amplified and 1p deletion negative NB and underwent chemotherapy according to the European unresectable NB protocol [19] before tumor surgical resection with minimal residual disease. Her younger brother, i.e. III-3, was diagnosed at age of 20 days with a \(\text{MYCN}\) gain and 1p deletion positive adrenal NB with multiple subcutaneous nodules and bone metastasis. He received chemotherapy according to the SIOP European NB Infant protocol trial 99.3 before underwent surgery on primary tumor with minimal residual disease. The first cousin III-1 had a stage 4S adrenal NB and laterocervical lymphonodes diagnosed at the age of 1 month. She underwent chemotherapy, surgery and radiotherapy on primary tumor bed [20]. All patients had good prognosis: III-1 and III-3 are currently in complete remission and III-2 is in minimal residual disease at 24 6/12 years, 3 10/12 years, 3 5/12 years from diagnosis, respectively.

Mutational screening of \(\text{PHOX2B}\) and \(\text{ALK}\) genes

Direct DNA sequencing analysis of \(\text{PHOX2B}\) and \(\text{ALK}\) genes was performed on the constitutional DNA from blood specimens of the three NB patients (Figure 1). No mutation was found in the three exons of \(\text{PHOX2B}\) (data not shown). Conversely, DNA sequences of exons 20 to 29 encoding the tyrosine kinase domain of \(\text{ALK}\) revealed a guanosine-to-cytosine change in exon 23, resulting in an arginine-to-proline substitution at codon 1192 (R1192P) in all three NB patients (Figure 1). DNA sequencing of their relatives indicated that the R1192P mutation was inherited from the unaffected mothers and the grandfather (II-1, II-2 and I-1, respectively; Figure 1). Since this \(\text{ALK}\) mutation was inherited from asymptomatic parents, additional genetic events are thought to be required for tumor development in these patients.

Array CGH analysis of somatic and germline DNA copy number changes in NB-affected siblings

To search for potential common chromosomal alterations indicating a putative predisposing NB locus, an array CGH analysis was performed on tumor and constitutional (peripheral blood) DNA of the affected siblings III-2 and III-3 (Figure 2). Array CGH analysis of tumor DNA revealed whole chromosome losses or gains in III-2, and partial chromosome imbalances in III-3 (Figure 2A, III-2 and III-3 depicted with green and red,
Figure 2. Array CGH copy number profiles of tumor and constitutional DNA samples from the affected siblings. Array CGH results from tumor (A) and constitutional (B) DNA analysis of the affected siblings (patient III-2 and III-3, in green and red, respectively). The X-axis represents the chromosomes, while the Y-axis represents the normalized log2 Cy5(patient)/Cy3(healthy control) fluorescence intensity thresholds -1 (loss) and 1 (gain), respectively. Array CGH results from tumor DNAs (A) revealed gains and losses of whole chromosome in stage 3 NB (patient III-2, in green) and partial chromosome imbalances in stage 4 NB (patient III-3, in red). Array CGH results from constitutional DNAs (B) revealed gains and losses of small chromosomal regions.

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respectively). In III-2, eight chromosomes (2, 6, 7, 13, 15, 17, 18
and 22) were entirely duplicated and four chromosomes (4, 8, 9
and X) were completely lost (Figure 2A). In III-3, the most
prominent copy number alterations affected the short arm of
chromosomes 1, 2 and 19 and the long arm of chromosomes 10,
14 and 17, where distinct gains and losses including loci for several
genes were detected (Figure 2A). However, no remarkable
alterations could orientate to any particular predisposing locus.

When array CGH analyses of the DNA tumor samples (Figure 2A)
were compared with matched constitutional DNA (Figure 2B),
three genomic gains were identified in regions devoid of copy
number variations (CNVs) at chromosomes 6q22, 14q24 and
15q15 that include loci for \( \text{PCNX} \) (pecanex homolog) and \( \text{CASC4} \) (cancer
susceptibility candidate 4), respectively (Figure 3 and Table 1).

These genomic gains are depicted in figures 3A, 3C and 3E for III-
2, while the gains at \( \text{PCNX} \) and \( \text{CASC4} \) were found in both
siblings (Figure 3C-F). To see whether these genomic gains were
shared with the affected first cousin III-1, array CGH analysis was
performed similarly on the constitutional DNA sample. In this NB
patient, none of these genomic gains were detected (Figure S1).

Next, to define which parent transmits these genomic alterations to
their offspring, array CGH analysis was performed also on the
case of both parents (Figure S1). The genomic gains in all
\( \text{NKAIN2} \), \( \text{PCNX} \) and \( \text{CASC4} \) loci were found in II-3 (Figure S2) who
does not carry the R1192P mutation of \( \text{ALK} \), but not in II-2 who
does carry the R1192 mutation.

\( \text{NKAIN2} \) is expressed at high levels in the affected siblings

To determine whether genomic gains detected by array CGH
resulted in enhanced gene expression \( \text{NKAIN2} \), \( \text{PCNX} \)
and \( \text{CASC4} \) mRNA levels were evaluated in the affected siblings
and their parents by qPCR. High \( \text{NKAIN2} \) expression (P<0.01) was
found in both individuals (II-3 and III-2) who carry the genomic
gain at \( \text{NKAIN2} \) as compared to healthy donors (Figure 4).

Surprisingly \( \text{NKAIN2} \) was expressed at high levels also in the
sibling III-3 who lacks the genomic gain, suggesting the
occurrence of additional mechanisms for \( \text{NKAIN2} \) gene regulation
(Figure 4). Conversely, genomic gains at \( \text{PCNX} \) and \( \text{CASC4} \) did not
result in enhanced gene expression, but rather in a
reduction. Hence, the latter two gains were not further
investigated.

\( \text{NKAIN2} \) is downregulated during neuronal differentiation
of NB cell lines

To investigate how \( \text{NKAIN2} \) expression may contribute to tumor
initiation and/or clinical phenotype, we next evaluated \( \text{NKAIN2} \)
mRNA levels on several NB cell lines without copy number
changes at 6p [21–23], in order to avoid the influence of somatic
DNA alterations on gene expression. The qPCR revealed that
\( \text{NKAIN2} \) was expressed at higher levels in the MYCN-amplified
NB cell lines [24], as compared to the less aggressive non-MYCN-
amplified NB cell lines (Figure 5A).

To determine whether \( \text{NKAIN2} \) is involved in NB cell
differentiation, two ATRAX-sensitive NB cell lines, SH-SY5Y
and SK-N-BE(2), were daily treated with ATRA for 4 and 10
days respectively, resulting in distinct morphological changes
with extensive neurite outgrowth. Differentiation was also
confirmed through the observed down-regulation of MYCN
and upregulation of NCAM at mRNA and protein levels,
respectively (Figure 5B). As shown in Figure 5B, ATRA
expression in both SH-SY5Y and SK-N-BE(2) cell lines,
suggesting a potential involvement of this gene in NB growth
and differentiation.

The coordinated expression of \( \text{NKAIN2} \) and MYCN indicates
the possibility that \( \text{NKAIN2} \) is regulated by MYCN. To address
the regulatory role of MYCN in the \( \text{NKAIN2} \) expression, we
utilized a MYCN-transfected NB cell line, Tet-21/N, carrying a
tetracycline-repressible MYCN transgene [25]. Tet-21/N cells
untreated or treated with doxycycline, were tested for the
expression of MYCN and \( \text{NKAIN2} \). MYCN expression was
drastically suppressed by doxycycline treatment in Tet-21/N, as
determined by qPCR and Western blotting (Figure 5C). However,
this clear change of MYCN expression, did not induce a
proportional change in the expression of \( \text{NKAIN2} \) (Figure 5C),
suggesting that MYCN is not directly involved in the regulation of
\( \text{NKAIN2} \) expression in NB cell lines.

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**Table 1. Array-CGH results in tumor and peripheral blood
dna samples.**

| Tumor DNA | Constitutional DNA |
|-----------|-------------------|
| **Cases** | **aCGH** | **Size (kb)** | **Genes** | **aCGH** | **Size (kb)** | **Genes** |
| III-2     | dup(6q22.31) | 333 | no genes | dup(6q22.31) | 333 | no genes |
|           | dup(6q22.31) | 156 | NKAIN2   | dup(6q22.31) | 156 | NKAIN2   |
|           | dup(14q24.2) | 258 | PCNX      | dup(14q24.2) | 258 | PCNX      |
|           | dup(15q15.3) | 90  | CASC4     | dup(15q15.3) | 90  | CASC4     |
| III-3     | del(1p36.33) | 19  | PCNX      | dup(14q24.2) | 258 | PCNX      |
|           | dup(2p25.3)  | 48  | CASC4     | dup(15q15.3) | 90  | CASC4     |
|           | del(10q22)   | 3   |           |               |     |           |
|           | dup(14q24.2) | 0.26|           |               |     |           |
|           | dup(17q21.2) | 41  |           |               |     |           |
|           | del(19p13.3) | 14  |           |               |     |           |
| II-2      | del(3p22.1)  | 145 | ULK4      |               |     |           |
|           | dup(1p36.23) | 37  | VAMP3     |               |     |           |
|           | dup(1p21.2)  | 24  | PER3      |               |     |           |
|           |               |     | VPS13D    |               |     |           |
| II-3      | dup(6q22.31) | 333 | no genes | dup(6q22.31) | 156 | NKAIN2   |
|           | dup(14q24.2) | 258 | PCNX      | dup(14q24.2)   | 258 | PCNX      |
| II-1      | del(2p16.3)  | 132 | no genes | dup(6q27)      | 453 | no genes |
|           | del(2p16.3)  | 132 | no genes | dup(6q27)      | 453 | no genes |

**Figure 3. Copy number alterations of chromosomes 6, 14 and 15 in the affected siblings.** The chromosome 6, 14 and 15 views (left) display the copy number profile of analysing constitutional DNA of III-2 (A, C and E) and III-3 (B, D and F) patients versus normal references. The gene views (right) magnify the selected amplification regions at 6q22, 14q24 and 15q15 indicating the log2 copy number ratios of individual oligos.

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Expression of NKAIN2 is significantly higher in the most aggressive NB lesions and the constitutional DNA of NB patients

Given the potential role of NKAIN2 in NB differentiation, expression of NKAIN2 was investigated in 12 primary NB lesions, including 5 low-risk and 7 high-risk NB, by immunohistochemistry. NKAIN2 was clearly detected on the cell membrane and cytoplasm of neuroblastic cells in 7/7 high-risk NB. Conversely, NKAIN2 was hardly seen in the neuroblastic cells of 5/5 low-risk NB. A representative staining revealing expression levels and cellular localization is shown in Figure 6A.

Finally, expression of NKAIN2 was investigated in the constitutional DNA of a cohort of 17 NB patients out of therapy by qPCR. As shown in Figure 6B, a significantly higher expression ($P = 0.002$) of $NKAIN2$ was observed in the blood of NB patients as compared to age-matched healthy donors. Taken together these data suggest that high NKAIN2 expression by either the gene amplification or some other mechanisms may be relevant to NB pathogenesis.

Discussion

Germline mutations in $ALK$ and $PHOX2B$ account for the majority of hereditary NBs. However, since these mutations are inherited and shared by unaffected parents, additional genetic events are thought to be required for tumor development.

To identify new candidate genes implicated in NB, we performed a comparison between somatic and germline DNA copy number changes in two NB-affected siblings carrying an $ALK$ germline activating mutation (R1192P). This ALK mutation has been already reported in familial NB [14]. Array CGH analysis revealed a germline genomic gain in $NKAIN2$ locus in one of the affected siblings, that was inherited from the parent who does not carry the $ALK$ mutation. Surprisingly, $NKAIN2$ was expressed at high levels also in the affected sibling that lacks the genomic gain at this locus, clearly suggesting mechanisms other than genomic gain involved in the gene regulation. We provided evidence that NKAIN2 play a role in NB development because its expression is directly related with the aggressiveness of human NB cell lines and tumor specimens and it is downregulated during ATRA differentiation. Herein, we distinguished two different NB cell phenotypes depending on the expression of NKAIN2: phenotype (a), neuroblastic cells that express high levels of NKAIN2; phenotype (b), neuroblastic cells with low NKAIN2 expression. The first phenotype was detected in the high-risk NB patients characterized by $MYCN$ amplification, poor prognosis and therapy resistance.

The second one features low-risk NB patients with low levels of MYCN associated with a relatively benign clinical profile, slow disease progression and therapy responsive. Furthermore, NKAIN2 was significantly expressed at higher levels in the peripheral blood of a cohort of NB patients as compared to age-matched healthy donors.

NKAIN2 belongs to a superfamily of transmembrane proteins interacting with $\beta$1 subunits of the Na+/K-ATPase encoded by four genes, $NKAIN1$, $NKAIN2$, $NKAIN3$ and $NKAIN4$ [26]. $NKAIN$ genes do not show any similarity with other known genes, although they share striking evolutionary conservation among species [26,27]. Amino acid conservation of the first two domains suggests that this protein family might function within membrane bilayer. Some authors showed that a Drosophila mutant carrying a P-element insertion in the first exon of Drosophila ortholog $dNKAIN$ displayed a temperature-sensitive paralysis [26]. The neuronal expression of $NKAIN$ proteins, their membrane localization and the phenotype of NKAIN Drosophila mutants strongly suggest that these proteins are critical for neuronal function. In agreement with this hypothesis, two patients with severe neurological impairment have been found to express a truncated form of $NKAIN2$ by means of a constitutional chromosomal translocation [27,28]. Because no other genes were disrupted, the clinical presentation of the two patients has been ascribed to $NKAIN2$ truncation [27,28]. As for $NKAIN2$ [26], mutations of Na+/K-ATPase alpha subunits in Drosophila have been also found to cause neural dysfunction, leading to seizures and neurodegeneration, suggesting their fundamental role in maintaining normal neuronal functions [29].

Although NKAIN2 does not seem to have any specific enzymatic activity, it could be involved in some essential cellular pathway as a regulator. Previous studies have indicated that different NKAIN2 transcripts are expressed in fetal and adult brain suggesting that they may play a specific role at different stages [27]. In situ hybridization experiments confirmed the tissue-specific expression of NKAIN2 in the central and peripheral nervous system, with a distribution consistent with expression in neurons [26].

Herein, we provide evidence that NKAIN2 is downregulated in two NB cell lines by ATRA treatment, suggesting a role in neuronal differentiation. Retinoic acid exerts essential effects during embryonic development and differentiation of several cell types [30] and has been studied as human anti-cancer drug as it induces cell proliferation arrest, morphological differentiation and apoptosis in several cancer types, including NB [31,32].

In summary, our data indicate that the $NKAIN2$ might contribute to the elucidation of the complex mechanism of NB differentiation.
and its genetic components. Additional functional studies around this Na/K-ATPase interacting gene are required to further investigate its role in the NB susceptibility and to unveil the molecular basis of NB development and progression, hopefully providing diagnostic and therapeutic advances in this field.

Figure 5. Expression of NKAIN2 in NB cell lines can be reduced by ATRA treatment. A, qPCR analysis of mRNAs from different NB cell lines with or without MYCN amplification (MNA). β-actin was used for normalization. B, qPCR analysis of MYCN and NKAIN2 mRNA expression (left panel) and immunoblot analysis of NCAM expression (right panel) in SH-SY5Y and SK-N-BE(2) cells untreated or treated with ATRA for 4 and 10 days, respectively. C, qPCR analysis of MYCN and NKAIN2 expression (left panel) and immunoblot analysis of MYCN expression (right panel) in Tet-21/N cells untreated or treated with doxycycline (Dox) for 5 days. The error bars represent the mean ± SD (n = 3).

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Materials and Methods

NB cell lines

All human NB cell lines, obtained from the American Type Culture Collection, were characterized by morphology and HLA class I typing by PCR-SSP (Genovision). Cell lines were maintained in RPMI 1640 medium (HTLA-230, SK-N-BE(2), SK-N-BE(2)c, and IMR-32), Dulbecco’s modified Eagle’s medium (SH-SY5Y), or Minimum Essential Medium Eagle (SK-N-SH). All media were supplemented with 10% heat-inactivated fetal bovine serum (HyClone), 2 mM glutamine, 100 units/ml penicillin and 100 μg/ml streptomycin. Tet-21/N cell line was kindly provided by Dr M. Schwab (Johann Wolfgang Goethe-University, Frankfurt, Germany, [25]). All-trans retinoic acid (ATRA) and doxycycline, both from Sigma-Aldrich, were used at 10 μM and 10 ng/L, respectively.

Ethics Statement

All tumor specimens and blood samples were obtained from NB patients diagnosed at the “Bambino Gesù” Children’s Hospital, Rome (Italy), after obtaining written informed parental consent and approval by the Ethical Committee of the Institution. The clinical information of NB patients including patients’ age at diagnosis, disease stage according to the International Neuroblastoma Staging System [18], MYCN oncogene copy number [33] and the International Neuroblastoma Pathology Classification [34] are listed in Table 2. They include 3 affected children of an Italian family of which two siblings and a first cousin, and 25 sporadic NB patients (Figure 1 and Table 2).

Genomic DNA extraction

Genomic DNA was extracted from peripheral blood and tumor samples with High Pure PCR Template Preparation Kit (Roche) according to the producer’s instructions. Concentration and purity

Figure 6. Expression of NKAIN2 in primary NB lesions and peripheral blood of NB patients. A. Immunohistochemistry of human NB tissues section with Ab to NKAIN2. Visualized with Fast Red (red), nuclei counter-stained with haematossilin (blue). NKAIN2 is strongly expressed on the cell membrane of neuroblasts in the high-risk NB (left panel) and undetectable in the neuroblasts of low-risk NB (right panel). Original magnification, ×40. Scale bars 30 μm. Data shown are representative of 7 high-risk and 5 low-risk NB. B, qPCR analysis of mRNA in 6 healthy donors (HD) and 17 NB patients in remission for at least 1 year (median age =6 for both HD and NB patients). β-actin was used for normalization. Significant differences between HD and NB patients were evaluated by Mann-Whitney test. doi:10.1371/journal.pone.0078481.g006
Table 2. Clinical-pathologic characteristics of NB patients.

| Characteristics category | Number (%) |
|--------------------------|------------|
| Age*                    |            |
| <18                     | 8 (29%)    |
| ≥18                     | 20 (71%)   |
| INSS Stage              |            |
| 1, 2, 3, 4S             | 17 (61%)   |
| 4                       | 11 (39%)   |
| MYCN                    |            |
| Amplified or gain       | 12 (43%)   |
| Not Amplified           | 13 (46%)   |
| Unknown                 | 3 (11%)    |
| Histology               |            |
| favorable               | 18 (64%)   |
| unfavorable             | 10 (36%)   |

INSS, International Neuroblastoma Staging System; histology is based on International Neuroblastoma Pathology Classification (INPC).

*Age at diagnosis; months.

DNA sequencing

Direct sequencing of the 3 exons of PHOX2B and of the exons 20 to 29 encoding the tyrosine kinase domain of ALK were performed using an Applied Biosystems 3500 Genetic Analyzer. Primers sequences and PCR conditions have been published elsewhere and are available [15,35]. Mutant fragments were confirmed by bi-directional and repeated sequencing, establishing new PCR reactions. Sequence data were analyzed via Mutation Surveyor v. 3.30 software (SoftGenetics).

Array-CGH and bioinformatic analysis

Array-CGH was carried out using a 60-mer oligonucleotide-based microarray platform that allows molecular profiling of genomic aberrations with an overall median probe spatial resolution of 13 Kb (180K) (Agilent Technologies Array-CGH Kits). Human genomic DNA was used as reference DNA. Aliquots of 2 µg of DNA from samples were double-digested with RsaI and AluI for 2 hours at 37°C. After heat inactivation of the enzymes at 65°C for 20 minutes, each digested sample was labelled by random priming (Agilent Technologies) for 2 hours using Cy3-dUTP for patient/parent DNAs and Cy5-dUTP for reference DNAs. Labelled products were column purified with Illustra CyScribe GFX purification kit (GE Healthcare). After probe denaturation and pre-annealing with 5-25 µg of Cot-1 DNA, hybridization was performed using an Applied Biosystems 7900-HT sequence detection system. Each sample was run in triplicate using pre-validated TaqMan gene expression assays from Applied Biosystems (Hs00900458_m1 for NKAIN2, Hs00900458_m1 for PCNX, Hs01036008_m1 for CASC4, Hs99999903_m1 for β-actin) under standard conditions. Relative gene expression was determined using the 2-Delta-Delta-Ct method [36], with β-actin as endogenous control.

Quantitative mRNA expression

Total RNA was extracted using Trizol Reagent (Invitrogen). First-strand cDNA was synthesized from 1 µg using the Super-Script II First Strand cDNA synthesis kit (Invitrogen). Quantitative real-time PCR (qPCR) reactions were performed using an Applied Biosystems 7900-HT sequence detection system. Each sample was run in triplicate using pre-validated TaqMan gene expression assays from Applied Biosystems (Hs009002833_m1 for NKAIN2, Hs00900458_m1 for PCNX, Hs01036008_m1 for CASC4, Hs99999903_m1 for β-actin) under standard conditions. Relative gene expression was determined using the 2-Delta-Delta-Ct method [36], with β-actin as endogenous control.

Antibodies, Immunoblotting and Immunohistochemistry

Whole-cell extracts obtained as previously described [24], were resolved on 8% SDS-PAGE and transferred onto Hybond-C paper (Amersham Bioscience). Filters were probed with anti-MYCN (sc-791, Santa Cruz Biotechnology), -NCAM1 (AB5032, Chemicon), or -β-actin (sc-1616, Santa Cruz Biotechnology) followed by incubation with peroxidase-coupled secondary Ab.

For immunohistochemical staining, tumor sections of paraffin-embedded tissue blocks were cut at 3 µm. Deparaffinization and antigen retrieval were performed with PT-link (Dako) in citrate buffer (pH 6.1) for 15 min at 98°C. Sections were incubated with the polyclonal rabbit anti-human NKAIN2 antibody (HPA045860, Sigma-Aldrich) for 45 min at room temperature followed by incubation with streptavidin alkaline phosphatase (Dako). Bound streptavidine was detected with Fast Red chromogene substrate (Dako) and levamisole in the reaction mixture for 10 min at room temperature. All samples were counterstained with haematoxylin. Sections of normal human placenta were used as positive control.

Statistical Analysis

Statistical significance was assessed by Mann-Whitney test. A P value of <0.05 was considered to be statistically significant.

Supporting Information

Figure S1 Copy number profiles of chromosomes 6, 14 and 15 in the affected cousin. The chromosome 6, 14 and 15 views (left) display the copy number profile of III-1 patient versus normal references. The gene views (right) magnify the selected amplification regions at 6q22, 14q24 and 15q15 indicating the log2 copy number ratios of individual oligos.

(TIF)

Figure S2 All copy number alterations in the affected siblings are inherited from the father. The chromosome views (left) display the copy number profile of constitutional DNA analysis from unaffected parents I-3 (A, C and E) and I-2 (B, D and F) of two affected siblings versus normal references. The gene views (right) magnify the selected amplification region at 6q22,
14q24 and 15q15 indicating the log2 copy number ratios of individual oligos.

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Author Contributions
Conceived and designed the experiments: PR, DF. Performed the experiments: PR CS MM LL AG PS. Analyzed the data: PR CS MM LL RB. Contributed reagents/materials/analysis tools: AG MAD AA FL RB. Wrote the paper: PR DF.