

Neurokinin-2 receptor polymorphism predicts lymph node metastasis in colorectal cancer patients

WEIJIA FANG^{1*}, CAIYUN FU^{2,3*}, XIAOGANG CHEN¹, XIAOZHOU MOU^{1,3}, FANLONG LIU¹,
JIONG QIAN¹, PENG ZHAO¹, YULONG ZHENG¹, YI ZHENG¹, JING DENG¹,
PINGJIANG YE⁴, YIFEI WANG⁵ and SHUSEN ZHENG¹

¹Biotherapy Center, Department of Medical Oncology, The First Affiliated Hospital, School of Medicine, Zhejiang University, Hangzhou, Zhejiang 310003; ²Lab of Proteomics and Molecular Enzymology, School of Life Sciences, Zhejiang Sci-Tech University, Hangzhou, Zhejiang 310018; ³Institute for Cell-based Drug Development of Zhejiang Province, Hangzhou, Zhejiang 310058; ⁴Department of Anorectum, People's Hospital of Shaoxing, Shaoxing, Zhejiang 312000; ⁵Second People's Hospital of Jiaxing, Jiaxing, Zhejiang 314000, P.R. China

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Abstract. To analyze the single nucleotide polymorphisms (SNPs) of two subtypes of neurokinin (NK) receptors, *NK1R* and *NK2R* (also known as *TAC1R* and *TAC2R*), in colorectal cancer (CRC), peripheral blood samples were collected from 199 CRC patients. Direct-sequencing was performed to identify the *NK1R* rs10198644 and *NK2R* rs4644560 SNPs. Genotype results were correlated with clinical factors. The allele frequencies of *NK1R* rs10198644 GC, CC and GG were 52, 17 and 31%, respectively, while that of *NK2R* rs4644560 GC, CC, and GG were 36, 50 and 14%, respectively. Patients with *NK2R* rs4644560 GC exhibited more positive lymph nodes than those with CC (mean, 2.2 vs. 1.3; $P=0.016$). Further analysis highlighted that the number of positive lymph nodes was also increased in the *NK2R* rs4644560 GC/*NK1R* rs10198644 GG group compared with the *NK2R* rs4644560 GG/*NK1R* rs10198644 GG group (mean, 2.2 vs. 0.9; $P=0.04$). These data suggested that the *NK2R* rs4644560 GC polymorphism alone or combination with *NK1R* rs10198644 GG may be a promising prognostic marker of lymph node metastasis in CRC patients.

Introduction

Neurokinins (NKs) are small neuropeptides that are present in the tumor microenvironment and can act on different stages of

carcinogenesis (1). Substance P (SP) and NK A (NKA) and B (NKB) are classical members of the NK family, which exert their biological effects by activating NK receptors denoted as *NK1R*, *NK2R* and *NK3R* (also known as *TAC1R*, *TAC2R* and *TAC3R*) (2). SP, the endogenous ligand for *NK1R*, activates responses correlated with tumor growth in several human cell lines bearing *NK1R* receptors (3). Moreover, NKA, the endogenous ligand for *NK2R*, is a good predictor of survival in patients with stage IV well-differentiated small bowel neuroendocrine neoplasms (4). Numerous tumors that express NKRs can misuse the NK-induced signaling of normal cells to promote the proliferation and survival of cancer cells, releasing cytokines and soluble mediators that favor tumor growth (5). However, the precise involvement and significance of NK in cancer pathologies remains to be fully defined.

NKRs belong to the family of seven transmembrane G-protein coupled receptors. *NK1R* and *NK2R* are dispersed throughout the central and peripheral nervous systems, while *NK3R* is distributed mainly in the central nervous system (6). Human *NK1R* and *NK2R* are 407- and 398-amino acid proteins, respectively (7). *NK2R* is expressed in a variety of organs, including the gastrointestinal tract, while functional data suggest that *NK2R* could be significant in mediating the NK-evoked smooth muscle contraction of organs (8). The prevalence of the *NK2R* mRNA α isoform indicates that *NK2R* is significantly involved in regulating human colonic functions (9).

In the United States, colorectal cancer (CRC) has the third highest incidence of new cases and the third highest rate of cancer-associated mortality, with 142,280 and 50,830 cases, respectively, in 2013 (10). It is also estimated that the mortality rate for CRC has been gradually increasing in China (11). The occurrence and development of this deadly disease may vary among patients; this may be attributed to variations in genotype patterns. Recently, a population-based epidemiological study in a Chinese population found that the *NK2R* rs4644560 genotype appeared to be associated with a decreased risk of CRC when combined with *NK1R* rs10198644 (12). This finding indicates that these two subtypes of *NKR* gene may

Correspondence to: Dr Shusen Zheng, Biotherapy Center, Department of Medical Oncology, The First Affiliated Hospital, School of Medicine, Zhejiang University, 79 Qinchun Road, Hangzhou, Zhejiang 310003, P.R. China
E-mail: shusenzheng@zju.edu.cn

*Contributed equally

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play an important role in CRC. The present study used blood samples collected from patients with CRC to clarify the potential association between the single nucleotide polymorphisms (SNP) of *NK2R* rs4644560/*NK1R*rs 10198644 and a range of clinical factors important in CRC.

Materials and methods

Participants. The present study was approved by the Ethics Committee of the First Affiliated Hospital, School of Medicine, Zhejiang University (Hangzhou, Zhejiang, China). Peripheral blood samples were collected from 199 patients with CRC following surgery in the First Affiliated Hospital, School of Medicine, Zhejiang University, between May and October 2013. All patients provided written informed consent.

Details of patient demographics and clinical information were recorded, including age, gender, alcohol consumption, tobacco exposure, primary tumor site, diagnostic stage, number of positive lymph nodes, differentiation grade and carcinoembryonic antigen (CEA) level. Lifestyle factors, including alcohol consumption and tobacco use, were defined as follows: An individual who consumed alcohol more than once per day for at least three months was considered as an alcohol drinker and an individual who had smoked more than one cigarette per day for over one year was considered as a tobacco user. Diagnostic stage was confirmed according to the National Comprehensive Cancer Network guidelines for CRC (13). Differentiation grade was defined as poor, moderate and well. An abnormal CEA level was defined as a level of >5 ng/ml.

Genotyping. Genomic DNA was extracted from whole blood samples using the QIAamp DNA Mini kit (Qiagen GmbH, Hilden, Germany) according to the manufacturer's instructions. Genotyping was performed with polymerase chain reaction (PCR) amplification and direct-sequencing. Primers were designed with Primer Premier 5.0 software (PREMIER Biosoft, Palo Alto, CA, USA) and synthesized by Genewiz Biotech (Beijing, China). The primer sequences used for amplification were as follows: *NK1R* forward, 5'-AAAGTTGGGATCTGCTTACACT-3' and reverse, 5'-TTTCTTTCTCTCACTGTCCCA-3'; and *NK2R* forward, 5'-ATGCTGCTGTGTCATCTGCT-3' and reverse, 5'-TATCTTGCC CAGGTTGGTCT-3'. The PCR was performed with 50 ng of genomic DNA, 0.25 μ M of each primer, 200 μ M of each deoxyribonucleotide, 3 mM MgCl₂ and 0.2 Units of Taq DNA polymerase (Takara, Dalian, China). The PCR amplification was performed as follows: Initial denaturation at 95°C for 5 min, followed by 9 cycles of denaturation at 95°C for 45 sec, annealing at 58°C for 45 sec (decreasing by 0.2°C every cycle) and elongation at 72°C for 30 sec; then 34 cycles of denaturation at 95°C for 45 sec, annealing at 55°C for 45 sec (decreasing by 0.2°C every cycle) and elongation at 72°C for 30 sec; and a final elongation at 72°C for 5 min.

The PCR products were purified using the SAP/EXO PCR kit (New England Biolabs, Inc., Ipswich, MA, USA). Two microliters of the purified amplicons were directly used for a sequencing reaction with the Big Dye Terminator cycle sequencing mix v3.1 (Applied Biosystems Life Technologies, Foster City, CA, USA). Dye purification was performed by alcohol/sodium acetate precipitation. Sequence analysis was

Table I. Patient characteristics (n=199).

Characteristic	Number (%)
Gender	
Male	117 (59)
Female	82 (41)
Location of primary tumor	
Left colon	34 (17)
Right colon	47 (24)
Rectum	118 (59)
Positive lymph nodes	
0-1	70 (35)
2-3	94 (47)
4-6	27 (14)
≥ 7	8 (4)
Alcohol drinker	
Yes	48 (24)
No	151 (76)
Tobacco user	
Yes	61 (31)
No	138 (69)

performed on an ABI 3730xl DNA analyzer (Applied Biosystems Life Technologies).

Statistical analysis. SPSS version 20.0 (SPSS, Inc., Chicago, IL, USA) was used for the analysis of the final results. $P < 0.05$ was considered to indicate a statistically significant difference. A two-tailed t-test (double samples heteroscedastic hypothesis) was used to clarify the association between the number of positive lymph nodes and tobacco exposure. Pearson's correlation was used to analyze the association between the number of positive lymph nodes and diagnostic stage. Bartlett's χ^2 test was used to analyze the association between the number of positive lymph nodes and the genotypes of each *NKR*. The data was assessed for homogeneity of variance following logarithmic transformation.

A one-way analysis of variance (ANOVA) was used to further clarify the impact of different genotype combinations of *NK2R* and *NK1R* on the number of positive lymph nodes. Prior to the analysis, the genotypes of *NK2R* and *NK1R* were first mixed into nine groups (Fig. 1).

Results

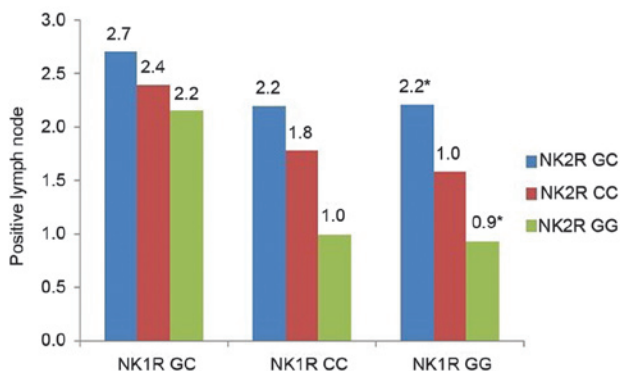
A total of 199 patients with pathologically confirmed CRC were enrolled in the study following surgical resection. The median age of the patients was 61 years (range, 23-84 years), and the detailed patient characteristics are listed in Table I. The allele frequencies of *NK1R* rs10198644 GC, CC and GG were 52, 17 and 31%, respectively, while for *NK2R* rs4644560 GC, CC and GG, the frequencies were 36, 50 and 14%. A comprehensive analysis of all patient factors, including demographic, lifestyle and clinical factors, was performed in order to clarify their association with the genotypes of *NK1R* and *NK2R*, as well as the interaction between them. The positive outcomes are described below.

Table II. Association between tobacco use and the number of positive lymph nodes.

Parameter	Tobacco	Non-tobacco	T-value	P-value (two-tailed)
Mean number of positive lymph nodes	3.2	2.0	2.10	0.04 ^a
Square deviation	16.5	3.1		
Number of patients	53	138		

^aP<0.05.Table III. Comparison of lymph nodes between the three genotypes of *NK2R* rs4644560.

Genotypes	Number of patients	Mean number of positive lymph nodes
GC	63	2.2 ^a
CC	79	1.3 ^a
GG	23	1.6
F-value		4.27
P-value		0.016 ^a

^aP<0.05.Figure 1. Comparison of median positive lymph node numbers within nine subtypes of *NK2R* and *NK1R* combinations. *P<0.05 *NK2R* GC/ *NK1R* GG vs. *NK2R* GG/ *NK1R* GG (P=0.04).

In total, there were 191 patients for whom information regarding positive lymph node numbers and tobacco exposure was available. Among these patients, 53 were in the tobacco group and 138 were in the non-tobacco group, and the mean number of positive lymph nodes was 3.2 and 2.0, respectively. The t-test showed that the tobacco group exhibited significantly more positive lymph nodes than the non-tobacco group (P=0.04; Table II).

In addition, among the 196 patients for whom information on stage and number of positive lymph nodes was available, Pearson's correlation found a positive correlation between the number of positive lymph nodes and the diagnostic stage of the tumor (r=0.48; P<0.001).

Finally, information pertaining to the number of positive lymph nodes and the *NKR* genotypes was available in 165 patients. Bartlett's χ^2 analysis showed that

NK2R rs4644560 GC was associated with significantly more positive lymph nodes than rs4644560 CC (mean, 2.2 vs. 1.3; P=0.016; Table III). Furthermore, the one-way ANOVA highlighted the interaction between *NK2R* rs4644560 and *NK1R* rs10198644, and showed that the number of positive lymph nodes was increased significantly in the rs4644560 GC/rs10198644 GG group compared with the rs4644560 GG/rs10198644 GG group (mean, 2.2 vs. 0.9; P=0.04), as shown in Fig. 1.

Discussion

Based on the identification of the role of NK pathway genes in the risk of CRC, the allele frequencies of SNPs in two selected subtypes of NK receptor, *NK1R* and *NK2R*, were investigated and their potential association with various clinical factors was analyzed. Notably, despite the limitation of the sample size, tobacco exposure, diagnostic stage and the selected *NKR* genotypes were all found to be significantly associated with the number of metastatic lymph nodes, which is well known to be one of the most important prognostic factors in CRC (14,15).

SNPs in receptor-encoded genes can affect a number of aspects of receptor function (16), and may also result in an increased susceptibility to disease or produce variable responses to therapeutic agents (17). There are several novel gene polymorphisms, including those for interleukin-10, SDF1 α and embryonic ectoderm development, which have been proven to be associated with lymph node metastasis in CRC (18,19).

It is well known that lifestyle factors, such as cigarette smoking and alcohol consumption, may contribute to the risk of cancer. In the present study, patients exposed to tobacco exhibited significantly more positive lymph nodes than those patients not exposed to tobacco; this result is consistent with the finding that tobacco exposure has a detrimental effect on

CRC (20). In agreement with a previously published study (21), the present study also found that diagnostic stage was associated with the number of metastatic lymph nodes.

As aforementioned, metastatic lymph nodes may be associated with a variety of clinical factors, therefore, its potential correlation with NK receptor polymorphisms, and their mutual interactions, was worthy of investigation. A previous study has shown that the stimulation of *NK2R* leads to partial blunting of the enhanced stimulatory effects mediated by *NK1R*. By contrast, stimulatory hematopoietic cytokines upregulate *NK1R* expression and downregulate the constitutively expressed *NK2R* in bone marrow stroma (22). In the present study, the role of *NK2R* rs4644560 GC in predicting lymph node metastasis was clarified based on the finding that this heterozygote is associated with more positive lymph nodes than the homozygote (CC) of *NK2R* rs4644560. Notably, despite the complexity of information on clinical interactions, the interactive model of *NK2R* and *NK1R* was also found to be useful. *NK2R* rs4644560 GC was associated with more positive lymph nodes than the corresponding GG genotype when combined with *NK1R* rs10198644 GG. Further research should be performed to elucidate the potential mechanism underlying this interaction.

Little has been known with regard to the mechanism of *NK2R* in CRC carcinogenesis. One study, however, began to shed light on this area. The study found that activation of *NK2R* in afferent neurons led to the protein kinase C (PKC)-induced phosphorylation of a certain gene (23). PKC δ was known to negatively modulate the canonical Wnt pathway and cell proliferation in colon tumor cell lines (24). Moreover, the connection between *NK2R* and PKC was confirmed by another study, which showed that stimulation of *NK2R* was not only linked to PKC activation, but that it was also associated with the activation of a multidrug resistance protein transporter *in vivo* (25). Extensive reviews have also described the neuropeptide receptors as targets for the treatment and diagnosis of cancer (5). Based on the present findings, one novel therapeutic strategy for colon cancer in the future may be the inhibition of the potential signaling pathways associated with certain NK receptor genotypes.

Pending further research to clarify the potential mechanism of the malignant inclination of the *NK2R* rs4644560 GC genotype or its combination with *NK1R* rs10198644 GG, the present study indicates that these genotypes may act as a promising prognostic marker, which may predict lymph node metastasis in CRC patients.

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