Association between a rare novel TP53 variant (rs78378222) and melanoma, squamous cell carcinoma of head and neck and lung cancer susceptibility in non-Hispanic Whites

Xiaoxiang Guan a, b, #, Li-E Wang b, #, Zhensheng Liu b, Erich M Sturgis b, c, Qingyi Wei b, *

a Department of Medical Oncology, Jinling Hospital, Medical School of Nanjing University, Nanjing, China
b Department of Epidemiology, The University of Texas MD Anderson Cancer Center, Houston, TX, USA
c Department of Head and Neck Surgery, The University of Texas MD Anderson Cancer Center, Houston, TX, USA

Received: December 12, 2012; Accepted: April 17, 2013

Abstract

Recently, several studies have investigated the association between a newly reported rare functional single nucleotide polymorphism (SNP) in TP53 (rs78378222) and cancer risk, but generated inconsistent findings. The present study further investigated this association with risk of melanoma, squamous cell carcinoma of head and neck (SCCHN) and lung cancer. Using volunteers of non-Hispanic Whites recruited for three large case–control studies, we genotyped the TP53 rs78378222 SNP in 1329 patients with melanoma, 1096 with SCCHN, 1013 with lung cancer and 3000 cancer-free controls. Overall, we did not observe any variant homozygotes in this study population, nor significant associations between the TP53 rs78378222AC genotype or C allele and risk for melanoma (P=0.680 and 0.682 respectively) and lung cancer (P=0.379 and 0.382 respectively), but a protection against SCCHN (P=0.008 and 0.008 respectively), compared with the AA genotype or A allele. An additional meta-analysis including 19,423 cancer patients and 54,050 controls did not support such a risk association either. Our studies did not provide statistical evidence of an association between this rare TP53 variant and increased risk of melanoma, nor of lung cancer, but a possible protection against SCCHN.

Keywords: biomarker ● genetic susceptibility ● genotype ● polymorphism

Introduction

Rare variants are more likely to have a functional impact and tend to have a greater effect size than do common variants [1, 2]. Thus, rare variants are likely to be a crucial genetic factor for human diseases, including cancer, and collectively they may contribute to a significant proportion of inherited susceptibility to cancer [3, 4]. The TP53 rs78378222, located in the 3′-untranslated region, is a newly identified rare SNP that is reportedly associated with risk of several cancers recently [5–7]. This rare variant is within the sole polyadenylation signal of TP53, with an A-to-C change in the sequence AATAAA to AATACA. This A-to-C sequence change may result in impairing proper termination and polyadenylation of the TP53 transcript and thus may alter cancer risk [6].

The first comprehensive and multi-institutional study identified a significant association between the TP53 rare rs78378222C allele and risk of prostate cancer, glioma and colorectal adenoma in Caucasian populations but not with risk of colorectal cancer, breast cancer and melanoma in the same study populations [6]. Subsequently, another two studies have reported that this variant allele is associated with significantly increased risk of oesophageal cancer in a Chinese population [7] and glioma in Caucasians [5]. In the present study, we tested the hypothesis that the TP53 rare rs78378222 SNP is associated with risk of aerodigestive tract cancers of the head and neck and lung, which share similar risk factors with oesophageal cancer, such as smoking, in our ongoing case–control studies of non-Hispanic Whites. In addition, we also tested the same hypothesis in our ongoing melanoma case–control study as well.

X. Guan and L. Wang contributed equally to this work.
*Correspondence to: Qingyi WEI, Department of Epidemiology, The University of Texas MD Anderson Cancer Center, Unit 1365, 1515 Holcombe Boulevard, Houston, TX 77030, USA. Tel.: 713-792-3020 Fax: 713-563-0999 E-mail: qwei@mdanderson.org

© 2013 The Authors. Journal of Cellular and Molecular Medicine Published by Foundation for Cellular and Molecular Medicine/Blackwell Publishing Ltd
This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.
doi: 10.1111/jcmm.12076
Materials and methods

Study population

Details of the recruitment of cases and controls have been reported elsewhere [8–10]. Briefly, the non-Hispanic White volunteers with histologically diagnosed melanoma (n = 1329) as well as SCCHN (n = 1096) and lung cancer (n = 1013), were recruited at The University of Texas MD Anderson Cancer Center between October 1999 and October 2007; the participation rate of eligible incident cases was approximately 95% of those who were initially contacted for participation. An additional 1926 cancer-free controls for both SCCHN and melanoma studies were recruited from among hospital visitors at MD Anderson Cancer Center during the same time period, and another 1074 cancer-free controls for lung cancer from the Kelsey-Seybold Clinics, Houston’s largest private multispecialty physician group. Cases and controls were frequency matched by age (±5 years), sex and ethnicity in each study. The study design, selection criteria, blood collection and DNA extraction have been described elsewhere [11].

TP53 rs78378222 SNP genotyping

Genotyping for the TP53 rs78378222 SNP was performed using the TaqMan assay with the Sequence Detection Software on an ABI Prism 7900 (Applied Biosystems, Foster City, CA, USA), the widely used genotyping platform. Primers and probes were supplied by Applied Biosystems. For all genotypes, the assay success rate was more than 99%, and the results of repeated assays for 10% of samples were 100% concordant. Associations between variant allele/genotypes and cancer risk were estimated by computing odds ratios (ORs) and their 95% confidence intervals (CIs) from both univariate and multivariable logistic regression models with or without adjustment for age, sex, smoking and drinking status. We performed the sequencing analysis for selected samples with different genotype as shown in Figure 1.

Mini meta-analysis

Because the association between TP53 rs78378222 SNP and cancer risk has been tested in several case–control studies with inconsistent results, we summarized published case–control association studies of TP53 variant rs78378222 and cancer risk, then conducted a meta-analysis for all published studies of Caucasians (excluded one Chinese study [7]), using the published data sets in addition to our data. All statistical methods were described elsewhere for association analysis [11] and for meta-analysis [12].

Statistical analysis

Differences in selected demographic variables, smoking status and alcohol use between cases and controls were evaluated by the chi-square test. The associations between TP53 rs78378222 SNP and cancer risk were estimated by computing ORs and their 95% CIs from both univariate and multivariable logistic regression models in case–control analysis. All tests were two-sided, and $P < 0.05$ was considered statistically significant. All statistical analyses were performed with SAS software (version 9.1.3; SAS Institute Inc., Cary, NC, USA), unless stated otherwise.

Results

Frequency distributions of age, sex, and TP53 rs78378222 between cancer patients and cancer-free controls

As shown in Table 1, there were no significant differences in distributions of age between controls and cases of SCCHN ($P = 0.722$), lung cancer ($P = 0.537$), melanoma ($P = 0.641$) and all cases combined ($P = 0.790$), nor in distributions of sex for SCCHN ($P = 0.537$), lung...
| Variables | SCCHN* | Lung cancer | Melanoma | Combined‡ |
|-----------|--------|-------------|----------|-----------|
|           | Cases  | Controls    |          | Cases     | Controls   |          | Cases     | Controls |
|           | n (%)  | n (%)       | p        | n (%)     | n (%)     | p        | n (%)     | n (%)    |
| Age (y)   |        |             |          |           |           |          |           |          |
| Range     |        |             |          |           |           |          |           |          |
| ≤50       | 300 (27.4) | 311 (28.6) | 0.722    | 210 (20.7) | 207 (19.2) | 0.567    | 611 (45.9) | 585 (44.6) | 0.641    | 1121 (32.6) | 988 (32.8) | 0.790 |
| 51–60     | 391 (35.6) | 389 (35.8) |          | 217 (21.4) | 247 (23.0) |          | 385 (28.9) | 401 (30.6) |          | 993 (28.8) | 847 (28.1) |          |
| >60       | 406 (37.0) | 386 (35.6) |          | 587 (57.9) | 622 (57.8) |          | 335 (25.2) | 325 (24.8) |          | 1328 (38.6) | 1180 (39.1) |          |
| Sex       |        |             |          |           |           |          |           |           |          |
| Male      | 826 (75.3) | 830 (76.4) | 0.537    | 457 (45.1) | 500 (46.5) | 0.521    | 790 (59.4) | 749 (57.1) | 0.247    | 2073 (60.2) | 1686 (55.9) | <0.01 |
| Female    | 271 (24.7) | 256 (23.6) |          | 557 (54.9) | 576 (53.5) |          | 541 (40.6) | 562 (42.9) |          | 1369 (39.8) | 1329 (44.1) |          |

*SCCHN: squamous cell carcinoma of head and neck.
†Chi-Square test.
‡Four hundred and fifty-eight controls overlapped between HN and melanoma studies.
| Studies       | Genotype* | Cases | Controls | MAF in Cases | MAF in Controls | Genotype p | Adjusted OR (95% CI)§ | Adjusted OR (95% CI)§ | Minimal detectable ORs with a 0.8 power |
|--------------|-----------|-------|----------|--------------|-----------------|------------|-----------------------|-----------------------|--------------------------------------|
| Melanoma     | AA        | 1292  |          |              |                 |            | 0.680                 | 1.15 (0.77–1.72)       | 0.682                  | 1.14 (0.77–1.71)                      |
|              | AC        | 37    |          | 0.01392      |                 | 0.680       | 1.15 (0.77–1.72)       | 0.682                  | 1.14 (0.77–1.71)                      |
| SCCHN        | AA        | 1083  | 3000     |              |                 |             |                       |                       | 0.455                  | 0.916                              |
|              | AC        | 13    | 77       | 0.00593      |                 | 0.008       | 0.44 (0.24–0.79)       | 0.008                  | 0.44 (0.24–0.80)                       |
| Lung Cancer  | AA        | 992   |          |              |                 |             |                       |                       | 0.441                  | 0.923                              |
|              | AC        | 21    |          | 0.01037      |                 | 0.379       | 0.84 (0.51–1.37)       | 0.382                  | 0.84 (0.51–1.37)                       |

* Nineteen samples failed in genotyping (one SCCHN, one lung cancer, two melanoma and 15 controls).
† No CC genotype was detected in this study population.
‡ Chi-Square test.
§ Logistic analysis, adjusted by age and sex.
SCCHN: squamous cell carcinoma of head and neck; MAF: minor allele frequency.
cancer ($P = 0.521$), melanoma ($P = 0.247$). Among the combined cases, 2073 (60.2%) were male, whereas 1686 (55.9%) were male in all the controls. The genotype frequencies of the TP53 rs78378222 SNP in both cases and controls were in agreement with the Hardy–Weinberg equilibrium.

Association between a rare novel TP53 variant (rs78378222) and cancer susceptibility in non-Hispanic Whites

As shown in Table 2, TP53 rs78378222 was not associated with overall cancer risk ($OR = 0.80; 95\% CI = 0.58–1.11$) in all cases combined, nor for risk of lung cancer ($OR = 0.85; 95\% CI = 0.48–1.53$) and melanoma ($OR = 1.10; 95\%; CI = 0.68–1.77$). However, this variant was protective for SCCHN risk ($OR = 0.41; 95\% CI = 0.21–0.79$), a finding apparently not to be consistent with those reported in the literature.

Mini meta-analysis on the association between the TP53 rs78378222C allele and overall cancer risk

We then conducted a mini meta-analysis of available data from published studies on the association between TP53 variant rs78378222 and cancer risk. Only three studies of Caucasian populations in addition to our data were included [5, 6], a Chinese study with small sample data set was excluded [7]. As a result, we found that, overall, the pooled data showed that TP53 rare variant was not significantly associated with cancer risk ($OR = 1.20; 95\% CI = 0.93–1.54$; Fig. 2).

Discussion

In this study of an association between the newly reported TP53 rs78378222 SNP (or C allele) and risk of melanoma as well as SCCHN and lung cancer in a non-Hispanic White population, we did not find a statistical evidence of an association between this rare TP53 SNP and increased risk of melanoma, nor for lung cancer. However, we did observe a protection against SCCHN risk, although this finding needs to be further validated in large studies.

In previously published studies, the newly reported TP53 variant rs78378222C allele was found to be associated with significantly increased risk of prostate cancer, glioma and colorectal adenoma, but not of colorectal cancer in Caucasian populations [6], and subsequent studies also confirmed the association for glioma in 566 cases and 603 controls in Caucasians [5] and reported an additional association with oesophageal cancer of the aerodigestive tract in 405 cases and 810 controls of a Chinese population [7].

In the present study, we intended to replicate the reported associations for melanoma and oesophageal cancer. A null association with melanoma risk is likely to be true, given the findings from our meta-analysis. However, our findings of no association with lung cancer but a protection against SCCHN are quite different from that of oesophageal cancer. Obviously, such an association is far from established, which may be explained by several possibilities. First, numerous mechanisms involved in dysfunction of p53 and its molecular network [13, 14], which are likely to have a more functional impact on carcinogenesis than this rare variant alone. Second, a spectrum of TP53 somatic mutations has likely participated in tumour promotion and progression that may be different in cancers of different origins [15]. Third, it is likely that the very same variant in the 3′-untranslated region of TP53 has a different role in somatic cells of different target tissues and in concert with different carcinogens exposed. Finally, there may be ethnic and geographic differences in the aetiology of dif-

Fig. 2 Meta-analysis of association between TP53 variant rs78378222 and cancer risk. OR and 95% CI were calculated using an allelic model for rs2736098. Guan X1-3 represented for SCCHN, lung cancer and melanoma studies, respectively; Stacey SN1-5 represented studies for prostate, glioma, colorectal, breast, melanoma and basal cell carcinoma cancers respectively.
different cancers, aerodigestive cancers in particular, in addition to cancer risk associated with this variant.

In our post-hoc power calculation (Table 2), we found that our sample size could detect minimal ORs of 0.489 or 1.666 for melanoma, 0.441 or 1.739 for lung cancer and 0.455 or 1.716 of SCCHN with a statistic power of 0.8, suggesting that our findings could be by chance, although the reported positive ORs ranged from 1.60 to 3.54 [5–7]. In conclusion, our results suggest that the novel but rare variant of TP53 is not associated with risk of melanoma, which is similar to the finding of a meta-analysis including 19,423 cancer patients and 54,050 controls; nor a risk association for lung cancer but a possible protection against SCCHN. However, larger and well-designed prospective studies are warranted to confirm our findings.

Acknowledgements

The authors thank Margaret Lung and Jessica Fiske for their assistance in recruiting the participants, Hongping Yu, Min Zhao and Jianzhong He, for their laboratory assistance. This work was supported by NIH grants R01 ES11740 and CA131274 (to Q. Wei), CA127219 and CA055769 (to M.R. Spitz), UT MD Anderson Cancer Center SPORE in Melanoma P50 CA093459, and P30 CA16672 (to MD Anderson Cancer Center).

Conflicts of interest

No potential conflicts of interest were disclosed.

References

1. Gorlov IP, Gorlova OY, Frazier ML, et al. Evolutionary evidence of the effect of rare variants on disease etiology. Clin Genet. 2011; 79: 199–206.
2. Lefevre JH, Bonilla C, Colas C, et al. Role of rare variants in undetermined multiple adenomatous polyposis and early-onset colorectal cancer. J Hum Genet. 2012; 57: 709–16.
3. Bodmer W, Bonilla C. Common and rare variants in multifactorial susceptibility to common diseases. Nat Genet. 2008; 40: 695–701.
4. Gorlov IP, Gorlova OY, Sunyaev SR, et al. Shifting paradigm of association studies: value of rare single-nucleotide polymorphisms. Am J Hum Genet. 2008; 82: 100–12.
5. Egan KM, Nabors LB, Olson JJ, et al. Rare TP53 genetic variant associated with glioma risk and outcome. J Med Genet. 2012; 49: 420–1.
6. Stacey SN, Sulem P, Jonassdotir A, et al. A germline variant in the TP53 polyadenylation signal confers cancer susceptibility. Nat Genet. 2011; 43: 1098–103.
7. Zhou L, Yuan Q, Yang M. A functional germ-line variant in the P53 polyadenylation signal and risk of esophageal squamous cell carcinoma. Gene. 2012; 506: 295–7.
8. Wei S, Wang LE, McHugh MK, et al. Genome-wide gene-environment interaction analysis for asbestos exposure in lung cancer susceptibility. Carcinogenesis. 2012; 33: 1531–7.
9. Liu Z, Ma H, Wei S, et al. Telomere length and TERT functional polymorphisms are not associated with risk of squamous cell carcinoma of the head and neck. Mol Carcinog. 2011; 50: 697–706.
10. Ma H, Zhou Z, Wei S, et al. Shortened telomere length is associated with increased risk of cancer: a meta-analysis. PLoS ONE. 2011; 6: e20466.
11. Reinhardt HC, Schumacher B. The p53 network: cellular and systemic DNA damage responses in aging and cancer. Trends Genet. 2012; 28: 128–36.
12. Sullivan KD, Gallant-Behm CL, Henry RE, et al. The p53 circuit board. Biochim Biophys Acta. 2012; 1825: 229–44.