Metabolic syndrome and the risk of urothelial carcinoma of the bladder: a case-control study

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Abstract

Background: The Metabolic syndrome (MetS) is an emerging condition worldwide, consistently associated with an increased risk of several cancers. Some information exists on urothelial carcinoma of the bladder (UCB) and MetS. This study aims at further evaluating the association between the MetS and UCB.

Methods: Between 2003 and 2014 in Italy, we conducted a hospital-based case-control study, enrolling 690 incident UCB patients and 665 cancer-free matched patients. The MetS was defined as the presence of at least three of the four selected indicators: abdominal obesity, hypercholesterolemia, hypertension, and diabetes. Odds ratios (ORs) and corresponding 95% confidence intervals (CIs) for MetS and its components were estimated through multiple logistic regression models, adjusting for potential confounders.

Results: Patients with MetS were at a 2-fold higher risk of UCB (95% CI: 1.38 – 3.19), compared to those without the MetS. In particular, ORs for bladder cancer were 2.20 (95% CI: 1.42 – 3.38) for diabetes, 0.88 (95% CI: 0.66 – 1.17) for hypertension, 1.16 (95% CI: 0.80 – 1.67) for hypercholesterolemia, and 1.63 (95% CI: 1.22 – 2.19) for abdominal obesity. No heterogeneity in risks emerged across strata of sex, age, education, geographical area, and smoking habits. Overall, 8.1% (95% CI: 3.9 – 12.4%) of UCB cases were attributable to the MetS.

Conclusions: This study supports a positive association between the MetS and bladder cancer risk.

Keywords: Bladder cancer, Diabetes, Metabolic syndrome, Obesity

Background

Bladder cancer ranks among the 10 highest incident cancers worldwide; it is one of the most frequent malignant tumours of the urinary system, with approximately 420,000 new cases each year among men and women, and a leading cause of cancer-related deaths [1, 2]. Incidence rates are three-to-four-fold higher in men than in women, and more than 90% of cases are urothelial carcinoma of the bladder (UCB). In Italy, standardized incidence rates for bladder cancer are 29.9 and 6.2/100,000 among men and women, respectively [3].

Tobacco smoking is a major risk factor for UCB, being responsible for 30% to 50% of cases in both sexes [4]. Other risk factors have been involved in UCB onset, including obesity, hypertension and diabetes [5–7]. The strong association with these medical conditions suggests a possible role of the metabolic syndrome (MetS) in UCB etiology [6, 7]. The MetS is a complex disorder described as a cluster of at least three risk factors for cardiovascular disease, including abdominal obesity, glucose intolerance, high blood pressure, high triglyceride levels and low high-density lipoprotein cholesterol levels [8].

The MetS has been consistently associated to increased risk for several cancers [6], of magnitude ranging from 1.1 to 1.6. Among women, the strongest associations were reported for endometrial, breast (postmenopausal), pancreatic and colorectal cancers [6, 7, 9, 10]. Among men, the strongest associations were with liver cancer, which persisted after adjustment for chronic infection with HBV/HCV [11], renal and colorectal cancer [6, 7, 11, 12].

Although the prevalence of the MetS is increasing worldwide and high rates of UCB are documented in
most countries, few epidemiological studies have been published on the relationship between the MetS and bladder cancer in the Mediterranean region. Two recent systematic reviews on the relationship between the MetS and UCB risk reported a positive association in men only [6, 7]. Therefore, to provide further information on the issue in a Mediterranean area, we examined data from an Italian case-control study investigating potential risk factors for UCB.

Methods
Between 2003 to 2014, we conducted a case-control study on urothelial carcinoma of the bladder within an established Italian network of collaborating centres, including Aviano and Milan in northern Italy, and Naples and Catania in southern Italy [13]. Cases were 690 patients aged 25 years or older (median age: 67 year; range: 25-84 years) with incident UCB admitted to major general hospitals in the study areas. Nearly all UCB (n = 642, 93.0 %) were confirmed by histological testing on tumour tissue specimen from biopsy or surgery. However, cases whose papillary features could not be determined (n = 138, 20 %) were excluded from the analysis of histological subtypes but were included in all other analyses. Overall, 268 UCB (38.8 %) were non-invasive (i.e., TNM pTis/Ta) and 307 (44.5 %) were well or modestly differentiated.

The control group included 690 patients frequency-matched to cases according to study centre, sex, and 5-year age group. Twenty-five controls were excluded after enrolment because of inappropriate admission diagnosis, thus leaving 665 eligible controls (median age: 66 years; range: 27-84 years). Controls were admitted to the same network of hospitals as cases for a wide spectrum of acute, non-neoplastic conditions unrelated to tobacco and alcohol consumption, to known risk factor for UCB, or to conditions associated to long-term diet modification. Overall, 28.9 % of controls were admitted for traumatic disorders, 22.1 % for non-traumatic orthopaedic disorders, 39.3 % for acute surgical conditions, and 9.8 % for other various illnesses. All study subjects signed an informed consent. Study protocol was approved by the Ethic Board of each study hospital (S. Maria degli Angeli hospital, register trial number 8/2004; and CRO Aviano National Cancer Institute, protocol number 590/D).

Trained interviewers administered a structured questionnaire to cases and controls during their hospital stay, thus keeping refusal below 5 % for both cases and controls. The structured questionnaire collected information on socio-demographic factors; lifetime smoking and alcohol drinking habits; dietary habits related to the two years preceding diagnosis/interview; problem-oriented medical history; and family history of cancer. Two specific sections investigated lifetime occupational exposure, and exposure to chemicals known (or suspected) to be related to UCB, including the use of hair dyes [13].

Information on clinical diagnosis of diabetes, drug-treated hypertension, and drug-treated hyperlipidaemia was self-reported and included age at diagnosis [14]. Diseases whose onset was less than one year before the interview were not considered. Likewise, self-reported height and weight one year prior to diagnosis/interview and at 30 and 50 years of age were collected.

Body mass index (BMI) was computed through the Quetelet’s formula (weight divided by squared height – kg/m²). The interviewers measured the waist circumference (2 cm above the umbilicus). The presence of abdominal obesity was defined using the International Diabetes Federation (IDF) cut-points (waist circumference ≥ 94 cm for men and ≥ 80 cm for women). Information on waist circumference could not be obtained for technical reason in 157 cases and 192 controls, thus leaving 533 cases and 473 controls for the present analysis shown in Tables 2 and 3. Sensitivity analyses were further conducted on all cases and controls using BMI ≥ 30 kg/m² as a proxy of abdominal obesity in patients missing waist circumference. MetS was determined according to the 2009 joint interim statement [15], as the presence of at least three of the following components: abdominal obesity, diabetes, drug-treated hypertension (as a proxy of elevated blood pressure), and drug-treated hyperlipidaemia (as a proxy of high triglyceride levels).

Odds ratios (ORs) and the corresponding 95 % confidence intervals (CIs) were calculated by means of unconditional logistic regression models, including terms for study centre, sex, 5-year age groups, years of education (i.e., <7, 7-11, ≥12) as a proxy of social status. To adjust for potential confounders (i.e. factors associated to both outcome and exposure), smoking habits (never; former; current: <20; 20+ cigarettes/day) were further included in the model. The test for trend was based on the likelihood-ratio test between the models with and without the linear term, reporting the median values in each strata of the variable of interest. Percent attributable risks (PAR) were computed using the distribution of risk factors among UCB cases [16].

Results
Most UCB cases were men and aged ≥65 years (Table 1). Cases and controls reported similar education, whereas current tobacco smoking was more frequent among UCB cases than controls (39.8 % and 21.7 %, respectively). Compared to never smokers, subjects smoking ≥20 cigarettes/day showed a seven-fold increased in UCB risk (95 % CI: 4.94-11.41), with a significant risk trend for number of cigarettes (P < 0.01).

Compared with people without any MetS components, the ORs were 2.00 (95 % CI: 1.17-3.41) for those with
three components and 7.93 (95 % CI: 1.71-36.79) for those with four components (P for trend < 0.01). After adjustment for the other MetS components, patients with diabetes (16.9 % cases and 8.0 % controls) showed a two-fold increase in UCB risk (95 % CI: 1.42-3.38 - Table 2). Likewise, patients reporting abdominal obesity showed a significantly higher UCB risk (OR = 1.63; 95 % CI: 1.22-2.19). No significant association emerged for treated hypertension (OR = 0.88; 95 % CI: 0.66-1.17) and treated hyperlipidaemia (OR = 1.16; 95 % CI: 0.80-1.67).

Compared to patients without indication of MetS (i.e., with two or less MetS indicators), those with MetS reported a two-fold increase in UCB risk (95 % CI: 1.38-3.19 - Table 2). Accordingly, in this study population, 8.1 % (95 % CI: 3.9-12.4 %) of all UCB cases were attributable to MetS (data not shown). Furthermore, 54 UCB cases (65.1 %) and 16 controls (38.1 %) with MetS reported diabetes; among these, the risk of UCB was 3.63 (95 % CI: 1.99-6.61) compared to those without MetS (Table 2). People with MetS without diabetes still showed a 16 % increased risk of UCB, but the association was not statistically significant.

The association between the MetS and UCB risk was similar in strata of gender (men vs. women; P for heterogeneity = 0.08), age (<65 years vs. ≥ 65 years; P = 0.59), education (<7 vs. ≥ 7 years; P = 0.92), geographical area (North vs. South of Italy; P = 0.43), and smoking habits (never, ever <20 cigarettes/day, and ever ≥ 20 cigarettes/day; P = 0.69 - Table 3). The association between the MetS and UCB risk was stronger (P = 0.03) for papillary UCB (OR = 2.61; 95 % CI: 1.68-4.04) than for non-papillary UCB (OR = 0.86; 95 % CI: 0.36-2.09), whereas no difference emerged according to tumour invasiveness (pTa/Tis vs. pT1-T4; P = 0.69 - Table 3). These results did not remarkably change using BMI ≥ 30 kg/m² when the information of waist circumference was missing (data not shown).

### Discussion

The present study supports a positive association between the MetS and risk of UCB, with a possible stronger association for papillary UCB. Conversely, no significant difference emerged according to gender, age, education, geographical area, and smoking habits. These findings are particularly interesting, giving the increasing prevalence of MetS worldwide and the attention by the scientific community on its effects on various health outcomes, including bladder and other urological cancers [6, 17].

In a prospective cohort study of 580 000 people – carried out within the Me-Can study – Haggstrom et al. [18] showed that MetS was associated with a significantly increased risk of UCB in men (RR = 1.10; 95 % CI: 1.01-1.18 for each incremental MetS unit), whereas no association was observed in women. Similarly, an Italian population-based study [19] observed a modest, non significant, increased risk of UCB only in men concurrently treated with antihyperglycaemic, antihypertensive, and hypolipidemic drugs. In their meta-analysis, Esposito et al. estimated that, in men, the presence of the MetS was significantly associated with the presence of UCB with a RR of 1.10 (95 % CI: 1.02-1.18) [11]. Our findings seem therefore to confirm
this association with a substantially stronger OR as compared to previous studies [6, 7].

Mechanisms that link MetS and cancer risk are not fully understood. However, some MetS components have been extensively investigated as cancer risk factors. There is mounting evidence showing the negative influence of obesity on genitourinary malignancies [20]. Several epidemiological studies showed a positive relationship between obesity and an increased risk of UCB, although others did not find any significant associations [21, 22]. Although BMI is generally used to define the grade of obesity, in our study we used abdominal obesity since it better explains obesity-related health risk [23]. Moreover a recent study used visceral obesity as individual component of MetS to predict adverse pathological features in UCB [24].

The biological mechanism for obesity-related carcinogenesis is not yet well characterized, but many possibilities have been suggested. High levels of adipose tissue correlate with high levels of cholesterol, a precursor for the androgen testosterone, which stimulates epithelial cell proliferation. High adipose levels have also been correlated with high plasma levels of vascular endothelial growth factor (VEGF), which both stimulate proliferation of epithelial cells. Adipose tissue also secretes leptin, which has been implicated in enhancing angiogenesis and, consequently, may also enhance tumour development [25]. Adiposity has also been associated with reduced mitochondrial function and, in turn, increased circulating reactive oxygen species, which can cause DNA damage [26].

The strongest single risk factor found in the present study was diabetes mellitus. Nonetheless, people with the MetS but without diabetes had a 16% increased risk of UCB. Furthermore, Table 2 shows a doubling of UCB risk in people with 4 MetS components (OR = 7.93; CI:

| Components | UCB Cases | Controls | OR (95% CI)\textsuperscript{a} | OR (95% CI)\textsuperscript{b} |
|------------|-----------|----------|-------------------------------|-------------------------------|
|            | n (%)     | n (%)    |                               |                               |
| Diabetes mellitus |          |          |                               |                               |
| No         | 443 (83.1) | 435 (92.0) | Ref                           | Ref                           |
| Yes        | 90 (16.9)  | 38 (8.0)  | 2.22 (1.45-3.39)              | 2.20 (1.42-3.38)              |
| Drug-treated hypertension |          |          |                               |                               |
| No         | 316 (59.3) | 279 (59.0) | Ref                           | Ref                           |
| Yes        | 217 (40.7) | 194 (41.0) | 0.99 (0.75-1.31)              | 0.88 (0.66-1.17)              |
| Drug-treated hyperlipidaemia |          |          |                               |                               |
| No         | 442 (82.9) | 399 (84.4) | Ref                           | Ref                           |
| Yes        | 91 (17.1)  | 74 (15.6)  | 1.21 (0.84-1.73)              | 1.16 (0.80-1.67)              |
| Abdominal obesity\textsuperscript{c} |          |          |                               |                               |
| No         | 145 (27.2) | 167 (35.3) | Ref                           | Ref                           |
| Yes        | 388 (72.8) | 306 (64.7) | 1.65 (1.23-2.21)              | 1.63 (1.22-2.19)              |
| Nr. of MetS components |          |          |                               |                               |
| None       | 83 (15.6)  | 85 (18.0)  | Ref                           |                               |
| 1          | 213 (40.0) | 208 (44.0) | 1.03 (0.70-1.51)              |                               |
| 2          | 154 (28.9) | 138 (29.2) | 1.23 (0.81-1.85)              |                               |
| 3          | 67 (12.6)  | 40 (8.5)   | 2.00 (1.17-3.41)              |                               |
| 4          | 16 (3.0)   | 2 (0.4)    | 7.93 (1.71-36.79)             |                               |
| \(\chi^2\) for trend; p-value |          |          |                               | 11.45; P < 0.01               |
| Increment of 1 MetS component |          |          |                               | 1.29 (1.11-1.49)              |
| Indicators of MetS\textsuperscript{d} |          |          |                               |                               |
| No         | 450 (84.4) | 431 (91.1) | Ref                           |                               |
| Yes        | 83 (15.6)  | 42 (8.9)   | 2.09 (1.38-3.19)              |                               |
| "Without diabetes" | 29 (5.4)  | 26 (5.5)   | 1.16 (0.65-2.07)              |                               |
| "With diabetes" | 54 (10.1) | 16 (3.4)   | 3.63 (1.99-6.61)              |                               |

\textsuperscript{a}Adjusted for sex, age (<55; 55-59; 60-64; 65-69; 70-74; ≥75 years), study centre, education (<7; 7-11; ≥12 years), and tobacco smoking (never; former; current: <20; 20+ cigarettes/day); \textsuperscript{b}Separate components were additionally adjusted for the other MetS components; \textsuperscript{c}According to IDF cut-points for waist circumference. \textsuperscript{d}Defined as the presence of at least three out of four MetS components.
1.71-36.79) than in people with diabetes (OR = 3.63; CI: 1.99-6.61) suggesting that the MetS indeed plays a relevant role in the risk of UCB.

Previous cohort studies have investigated the association between diabetes and UCB risk [27–31]. All these studies reported an increased UCB risk for both men and women with diabetes levels, but strongest associations were seen with longer diabetes duration [27]. The Me-Can study reported a statistically significant increased risk among women, with an RR of 1.45 (95 % CI: 1.05–2.01) per mmol increment of glucose [32]. A possible additional pathway between diabetes and UCB risk is the increased incidence of urinary tract infections among subjects with diabetes [30]. Among separate components of MetS, high blood pressure and hypercholesterolemia were not significantly associated with the risk of UCB in our study. These findings are consistent with those of previous prospective studies [6, 18].

Possible study limitations included selection and information bias. The proportion of pTa/Tis in our case series (45.3 %) is slightly lower than expected (approximately 60 %), thus limiting the generalization of our results. However, similar associations were found for pTa/Tis and pT1-T4 UCB, suggesting that this type of selection bias had a limited impact on our results. Information on MetS components was based on self-reported data from a questionnaire, which collected history of diabetes, treated hypertension, and treated hyperlipidaemia, rather than direct measurements of fasting plasma glucose, blood pressure, triglycerides and HDL cholesterol. Underestimation of the prevalence of MetS may therefore have occurred. However, reliability of our questionnaire on diabetes was tested among almost 300 subjects who were interviewed twice, reporting a satisfactory agreement (k statistic = 0.85) [14]. Moreover, a recent cohort study from Spain showed that self-reported data on MetS indicators and on MetS itself are sufficiently accurate for epidemiological inference [32]. Likewise, validation studies of hypertension confirmed with a medical examination found a reasonable accuracy of self-reported information

Table 3 Odds ratio (OR) and corresponding 95 % confidence interval (CI) for urothelial carcinoma of the bladder (UCB), according to indicators of the metabolic syndrome (MetS) in strata of selected variables. Italy, 2003-2014

| Variables | Indicator of MetS | OR (95 % CI) | \( \chi^2 \) for heterogeneity; \( p \)-value |
|-----------|-------------------|-------------|--------------------------------------|
|           | No | Ca:Co | Yes | Ca:Co |           |            |
| Sex       |     |       |     |       |           |            |
| Men       | 381:359 | 75:32 | 2.49 (1.56-3.96) |            |            |
| Women     | 69:72   | 8:10  | 0.81 (0.25-2.60) | 3.08; \( p = 0.08 \) |            |
| Age (years) |     |       |     |       |           |            |
| <65       | 183:199 | 25:14 | 2.53 (1.21-5.20) |            |            |
| ≥65       | 267:232 | 58:28 | 1.97 (1.18-3.29) | 0.31; \( p = 0.59 \) |            |
| Education (years) |     |       |     |       |           |            |
| <7        | 178:185 | 43:24 | 2.07 (1.14-3.75) |            |            |
| ≥7        | 271:246 | 40:18 | 2.16 (1.17-3.97) | 0.01; \( p = 0.92 \) |            |
| Geographical area |     |       |     |       |           |            |
| North     | 307:311 | 64:31 | 2.31 (1.42-3.74) |            |            |
| South     | 143:120 | 19:11 | 1.56 (0.66-3.67) | 0.61; \( p = 0.43 \) |            |
| Smoking habit |     |       |     |       |           |            |
| Never     | 67:150   | 7:16  | 1.16 (0.43-3.14) |            |            |
| Ever <20 cig./day | 190:174 | 34:13 | 2.45 (1.23-4.91) |            |            |
| Ever ≥20 cig./day | 186:104 | 41:13 | 1.85 (0.93-3.70) | 1.37; \( p = 0.57 \) |            |
| Histological subtype |     |       |     |       |           |            |
| Non-papillary | 82:431 | 7:42  | 0.86 (0.36-2.09) |            |            |
| Papillary  | 299:431 | 69:42 | 2.61 (1.68-4.04) | 4.82; \( p = 0.03 \) |            |
| Invasiveness |     |       |     |       |           |            |
| pTa/Tis   | 183:431 | 41:42 | 2.32 (1.42-3.79) |            |            |
| pT1-T4    | 221:431 | 38:42 | 2.13 (1.28-3.57) | 0.16; \( p = 0.69 \) |            |

\( a \)Adjusted for sex, age (<55; 55-59; 60-64; 65-69; 70-74; ≥75 years), study centre, education (<7; 7-11; ≥12 years), and tobacco smoking (never; former; current; <20; ≥20+ cigarettes/day);

\( b \)The sum does not add up to the total because of missing values.
Other self-reported MetS indicators might have been underestimated, but such information bias is likely to have occurred similarly in cases and controls, and, consequently, should have led to an attenuation of the real association [16, 34].

Other potential limitations of this study design comprise recall bias, since cases might have been more sensitized than controls to report history of disease. The hospital setting should have, however, improved information comparability of cases to controls, because both groups were interviewed under similar conditions and were therefore similarly sensitized toward recalling medical history. However, cases and controls were enrolled from the same catchment areas, and careful attention was paid to exclude from the control group subjects admitted for any condition related to the exposures under study, including tobacco smoking. Furthermore, results were consistent when analyses were performed excluding, in turn, the main diagnostic categories of controls. On the other hand, our findings were strengthened by the nearly complete participation of identified cases and controls, and by the use of a validated, reproducible questionnaire [14].

Conclusion
Our data suggest that metabolic aberrations related to the MetS, which are known to increase the risk of several types of cancer, also increase the risk of UCB. Both the worldwide increasing MetS prevalence and the rising incidence of UCB suggest that each year a considerable fraction of this cancer is attributable to the MetS. Thus, evidence is needed to investigate whether effective interventions to reduce the prevalence of the MetS in adult populations could reduce UCB risk. Moreover, patients with the MetS, even in presence of obesity and/or diabetes, should be encouraged to follow appropriate cancer control strategies.

Abbreviations
BMI: Body mass index; Ca:Co: Cases:Controls; CI: Confidence interval; Cig: Cigarettes; HBV: Hepatitis b virus; HCV: Hepatitis c virus; IDF: International Diabetes Federation; MetS: Metabolic syndrome; OR: Odds ratio; PAR: Percent attributable risk; UCB: Urothelial carcinoma of the bladder; VEGF: Vascular endothelial growth factor.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
MM, JP and CLV study concepts, MM, JP and DS study design, MG, RT and ML data acquisition, MDM, PC, AG, CB and MM data analysis and interpretation, AC, MDM and FT statistical analysis, MM, JP and MDM manuscript preparation, MDM, AC and CB manuscript editing, GC, CLV and DS manuscript review. All authors read and approved the final manuscript.

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References
1. Mutma-Nascimento C, Schmitz-Dager BJ, Zeegers MP, Steineck G, Kogevinas M, Real FX, et al. Epidemiology of urinary bladder cancer: from tumor development to patient’s death. World J Urol. 2007;25:285–95.
2. Babjuk M, Burger M, Zigeuner R, Shahid SF, Van Rhijn BW, Compérat E, et al. European Association of Urology. EAU guidelines on non-muscle-invasive urothelial carcinoma of the bladder: update 2013. Eur Urol. 2013;64:639–53.
3. Forman D, Bray F, Brewster DH, Gombe Mbilavice C, Kohler B, Pires M, et al. Cancer Incidence in Five Continents, Vol. X. IARC Sci Pub, Lyon, 2013.
4. Freedman ND, Silverman DT, Hollenbeck AR, Schatzkin A, Abnet CC. Association between smoking and risk of bladder cancer among men and women. JAMA. 2011;306:737–45.
5. Qin Q, Xu X, Wang X, Zheng XY. Obesity and risk of bladder cancer: a meta-analysis of cohort studies. Asian Pac J Cancer Prev. 2013;14:3117–21.
6. Stocks T, Bjorge T, Uliman H, Manjer J, Häggström C, Nagel G, et al. Metabolic risk score and cancer risk: pooled analysis of seven cohorts. Int J Epidemiol. 2015;44:1353-63.
7. Cantello F, Ciccone A, Salonia A, Autonimo R, De Nunzio C, Briganti A, et al. Association between metabolic syndrome, obesity, diabetes mellitus and oncological outcomes of bladder cancer: A systematic review. Int J Urol. 2015;22(2):32–
8. Kassi E, Pervanidou P, Kaltsas G, Chrousos G. Metabolic syndrome: definition and controversies. BMC Med. 2011;9:48.
9. Esposito K, Capuano A, Giugliano D. Metabolic Syndrome and cancer: holistic or reductionist? Endocrine. 2014;45:362–4.
10. Rosato V, Zucchetto A, Bosetti C, Dal Maso L, Montella M, Pelucchi C, et al. Metabolic syndrome and endometrial cancer risk. Ann Oncol. 2011;22:884-9.
11. Esposito K, Chioldi P, Colao A, Lenzi A, Giugliano D. Metabolic syndrome and risk of diabetes. Care. 2012:35:2402–11.
12. Turati F, Talimani R, Pelucchi C, Polese J, Franceschi S, Crispo A, et al. Metabolic syndrome and hepatocellular carcinoma risk. Br J Cancer. 2013;108:222–8.
13. Polese J, Bosetti C, Di Maso M, Montella M, Libra M, Garbiglio A, et al. Tobacco smoking and the risk of papillary and non-papillary transitional cell carcinoma of the bladder. Cancer Causes Control. 2014;25:1151-8.
14. Bosetti C, Tavani A, Negri E, Trichopoulou D, La Vecchia C. Reliability of data on medical conditions, menstrual and reproductive history provided by hospital controls. J Clin Epidemiol. 2001;54:902–6.
15. Alberti KG, Eckel RH, Grundy SM, Zimmet PZ, Cleeman JI, Donato KA, et al. Harmonizing the metabolic syndrome: a joint interim statement of the International Diabetes Federation task force on epidemiology and prevention; National Heart, Lung, and Blood Institute; American Heart Association; World Heart Association; International Atherosclerosis Society; and International Association for the Study of Obesity. Circulation. 2009;120:1640–45.

16. Breslow NE, Day NE. Statistical methods in cancer research, Vol. 1. The analysis of case-control studies. IARC Sci Publ. 1980;32:5–338.

17. Pelucchi C, Serraino D, Negri E, Montella M, Dellanone C, Talamini R, et al. The metabolic syndrome and risk of prostate cancer in Italy. Ann Epidemiol. 2011;21:395–41.

18. Haggstrom C, Stocks T, Rapp K, Bjärge T, Lindkvist B, Concinni H, et al. Metabolic syndrome and risk of bladder cancer; prospective cohort study in the metabolic syndrome and cancer project (Me-can). Int J Cancer. 2011;128:1890–8.

19. Russo A, Autelitano M, Bisanti L. Metabolic syndrome and cancer risk. Eur J Cancer. 2008;44:293–7.

20. Jansson L, Katzmarzyk PT, Ross R, Waist circumference and not body mass index explains obesity-related health risk. Am J Clin Nutr. 2004;79:79–84.

21. Holick CN, Giovannucci EL, Stampfer MJ, Michaud DS. Prospective study of body mass index, height, physical activity and incidence of bladder cancer in US men and women. Int J Cancer. 2007;120:40–6.

22. Koebnick C, Michaud D, Moore SC, Park Y, Hollenbeck A, Ballard-Barbash R, et al. Body mass index, physical activity, and bladder cancer in a large prospective study. Cancer Epidemiol Biomarkers Prev. 2008;17:1214–21.

23. Matsushita Y, Nakagawa T, Shiratori M, Yamamoto S, Takahashi I, Mizoue T, et al. How can waist circumference predict the body composition? Diabetol Metab Syndr. 2014;6:11.

24. Cantelli F, Ciccone A, Autononi R, Salonia A, Briganti A, Ferro M, et al. Visceral obesity predicts adverse pathological features in urothelial bladder cancer patients undergoing radical cystectomy: a retrospective cohort study. World J Urol. 2014;32:559–64.

25. Cirillo D, Raghiglio AM, La Montagna R, Giordano A, Normanno N. Leptin signaling in breast cancer: an overview. J Cell Biochem. 2008;105:956–64.

26. Beckman KB, Ames BN. Oxidative decay of DNA. J Biol Chem. 1997;272:19633–6.

27. Newton CC, Capetanakis MT, Campbell PT, Jacobs EJ. Type 2 diabetes mellitus, insulin-use and risk of bladder cancer in a large cohort study. Int J Cancer. 2013;132:896–91.

28. Larsson SC, Orsini N, Brismar K, Wolk A. Diabetes mellitus and risk of bladder cancer: a meta-analysis. Diabetologia. 2006;49:289–293.

29. Tseng CH. Diabetes and risk of bladder cancer: a study using the National Health Insurance database in Taiwan. Diabetologia. 2011;54:2099–102.

30. Mackenzie T, Zemski MS, Ferrara A, Schned A, Karagas MR. Diabetes and risk of bladder cancer: evidence from a case-control study in New England. Cancer. 2011;117:1525–32.

31. Prizment AE, Anderson KE, Yuan JM, Folsom AR. Diabetes and risk of bladder cancer among postmenopausal women in the Iowa Women’s Health Study. Cancer Causes Control. 2013;24:603–8.

32. Stocks T, Rapp K, Bjärge T, Manjer J, Ulmer H, Selmer R, et al. Blood glucose and risk of incident and fatal cancer in the metabolic syndrome and cancer project (me-can): analysis of six prospective cohorts. PLoS Med. 2009;6:e1000201.

33. Barrio-Lopez MT, Ben-Rastrollo M, Bruzzi PJ, Fernandez-Montero A, Garcia-Lopez M, Martinez-Gonzalez MA. Validation of metabolic syndrome using medical records in the SUN cohort. BMC Public Health. 2011;11:867.

34. Vargas CM, Burt VL, Gillum RF, Pamuk ER. Validity of self-reported hypertension in the National Health and Nutrition Examination Survey III, 1988–1991. Prev Med. 1997;26:678–85.