An etiologic prediction model incorporating biomarkers to predict the bladder cancer risk associated with occupational exposure to aromatic amines: a pilot study

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Abstract

Background: No etiological prediction model incorporating biomarkers is available to predict bladder cancer risk associated with occupational exposure to aromatic amines.

Methods: Cases were 199 bladder cancer patients. Clinical, laboratory and genetic data were predictors in logistic regression models (full and short) in which the dependent variable was 1 for 15 patients with aromatic amines related bladder cancer and 0 otherwise. The receiver operating characteristics approach was adopted; the area under the curve was used to evaluate discriminatory ability of models.

Results: Area under the curve was 0.93 for the full model (including age, smoking and coffee habits, DNA adducts, 12 genotypes) and 0.86 for the short model (including smoking, DNA adducts, 3 genotypes). Using the "best cut-off" of predicted probability of a positive outcome, percentage of cases correctly classified was 92% (full model) against 75% (short model). Cancers classified as "positive outcome" are those to be referred for evaluation by an occupational physician for etiological diagnosis; these patients were 28 (full model) or 60 (short model). Using 3 genotypes instead of 12 can double the number of patients with suspect of aromatic amine related cancer, thus increasing costs of etiologic appraisal.

Conclusions: Integrating clinical, laboratory and genetic factors, we developed the first etiologic prediction model for aromatic amine related bladder cancer. Discriminatory ability was excellent, particularly for the full model, allowing individualized predictions. Validation of our model in external populations is essential for practical use in the clinical setting.

Keywords: Urinary bladder neoplasms, Occupational exposure, Risk, Logistic models, ROC curve

Background

Bladder cancer (BC) accounts for 5–10% of all malignancies among males in Europe and USA [1]. The most important risk factors are smoking, genetic susceptibility and occupational exposure [2].

An excess BC risk was identified since the early 1950s in the rubber industry and was associated to the use of b-naphthylamine [3]. Small excesses of BC risk have continued to be observed even in more recent studies of rubber workers published in the 1980s and 1990s [4]. Other aromatic amines (AAs) have been shown to be carcinogenic [5]. Nonetheless, occupational exposure to AAs has continued due to their industrial and commercial value [6]. AAs exposure can occur at lower extent in many other occupational settings. A systematic review of Italian studies estimated that 4 to 24% of BCs are attributable to occupational exposure [7]. BC cases effectively compensated by INAIL were 440 from 2000 to 2006, on average 63 per year [8]. Because in Italy incident cases of BC in 2006 were about 17,000/year in males [9], the expected cases of occupational exposure...
disease should be between 680 (17,000×0.04) and 4080 (17,000×0.24), and underreporting between 91% ((63/680) -1) and 98% ((63/4080) -1).

There is a strong genotoxic mechanism for carcinogenicity of several AAs, and multiple metabolic pathways as well as many polymorphic genes have been found to be implicated in the activation of AAs into DNA-reactive intermediates [10]. Over the last 20 years, numerous biomarkers have been investigated in workers exposed to AAs [6]. A key biomarker are the DNA adducts which are considered as the ‘biologically effective dose’ because they represent an integrated measure of carcinogen exposure, absorption, distribution, metabolism and DNA repair [11].

Few studies have combined clinical factors with blood and urinary biomarkers into risk profiles that can be used to predict the likelihood of etiological diagnosis of BC [12–16]. To the best of our knowledge, no study has tried to use biomarkers to predict the etiological diagnosis of occupational BC. The causal attribution to occupation usually relies on thorough occupational history collection, pertinent and documented risk assessment, availability of industrial hygiene measurements, appraisal and reference to evidence based literature data; to reach etiological diagnosis, the likely best option is to refer patients to occupational health specialists. An etiologic prediction model tool divides an initial population of BC cases into a smaller fraction of ‘positives’ that should be referred to an occupational health specialist for etiologic assessments, and a larger portion of ‘negatives’ that should no longer be considered. To be effective, an algorithm should increase the number of single BC cases into a smaller fraction of cases; the local Ethical Committee approved the study.

Methods
Study design and population
The present study includes the “cases” arm stemming from of an earlier hospital-based case-control study fully described in previous publications [17–20]. Inclusion criteria were being male, aged 20–80, Italian. Cases were all 199 newly diagnosed, histologically confirmed BC patients, admitted to the Urology Departments of two large hospitals from 1997 to 2000. Controls were all 214 non-neoplastic urological patients matched to cases by age (±5 years), period and hospital of admission. A written informed consent was obtained from each subject; the local Ethical Committee approved the study.

Data collection
Peripheral blood lymphocytes (PBLs) were collected and automated DNA extraction was performed according to Extragen kit [Extragen BC by TALENT] [17]. Genotyping of glutathione S-transferase M1 (GSTM1) null, GSTT1 null, GSTP1 I105V, N-acetyltransferase 1 (NAT1) fast, NAT2 slow, cytochrome P450 1B1 (CYP1B1) V432 L, sulforaphase 1A1 (SULT1A1) R213H, myeloperoxidase (MPO) G-463A, catechol-O-methyltransferase (COMT) V108 M, manganese superoxide dismutase (MnSOD) A-9 V, NAD(P)H:quinone oxidoreductase (NQO1) P187S, X-ray repair cross-complementing group 1 (XRCC1) R399Q, XRCC3 T241 M, and xeroderma pigmentosum complementation group (XPD) K751Q polymorphisms was assessed using Amplification Refractory Mutation System assay [17]. Bulky-DNA adducts were detected by 32P–postlabeling after Nuclease P1 enrichment and labelled adducts resolution on Thin Layer Chromatography (TLC) [20]. DNA adduct levels were measured as relative adduct level per 10⁶ nucleotides. A trained interviewer collected information on demographic variables, lifetime smoking history, coffee and other liquid consumption, dietary habits, lifetime occupation history by questionnaire. Job titles and individual activities, as well as occupational exposures to AAs, were blindly coded by an occupational physician according to methodology described in previous publications [21]. Occupations involving exposure to AAs were attributed to 11 International Standard Classification of Occupations (ISCO, International Labour Office, 1968) codes for job tasks (1–61.30: Painter, Artist; 3–70.20: Mail Sorting Clerk; 5–70.30: Barber-Hairdresser; 7–41.40: Mixing- and Blending-Machine Operator, Chemical and Related Processes; 8–01.10: Shoemaker, General; 8–11.20: Cabinetmaker; 8–73.70: Vehicle Sheet-Metal Worker; 9–01.35: Rubber Moulding-Press Operator; 9–31.20: Building Painter; 9–39.20: Brush-Painter, except Construction; 9–39.30: Spray-Painter, except Construction) and 11 International Standard Industrial Classification of all Economic Activities (ISIC, United Nations, 1968) codes for industrial activities (3240: Manufacture of footwear; except vulcanized or moulded rubber or plastic footwear; 3320: Manufacture of furniture and fixtures, except primarily of metal; 3521: Manufacture of paints, varnishes and lacquers; 3559: Manufacture of rubber products not elsewhere classified; 3819: Manufacture of fabricated metal products except machinery and equipment not elsewhere classified; 3824: Manufacture of special industrial machinery and equipment except metal- and wood-working machinery; 3843: Manufacture of motor vehicles; 5000: Construction; 9415: Authors,
music composers and other independent artists not elsewhere classified; 9513: Repair of motor vehicles and motorcycles; 9591: Barber and beauty shops).

**Best-case definition**

We investigated exposure to AA in all jobs held during lifetime, carefully assessing the level and the temporal aspects of such exposure according to standardized procedures [21]. Then, in order to achieve an optimal case definition [22], the critical values for time since first exposure (TSFE) and time since last exposure (TSLE) were chosen based on literature findings. In a cohort of Italian dyestuff workers [23], the risk of BC mortality decreased with increasing TSLE and became non-significant at ≥30 years since last exposure. Out of 19 BC patients observed in a Japanese dyestuff-plant, 17 showed a TSFE ≥20 years and 18 a TSLE ≤35 years [24]. Thus, the criteria for best-case definition were: TSFE higher than 20 years; TSLE lower than 35 years; length of exposure of at least 1 year; any value of cumulative exposure to AAs. The 15 BC cases complying with the above criteria were considered AA-related bladder cancer.

**Variables and statistical analyses**

Smoking was a categorical variable with three levels: nonsmokers; former smokers from >20 years; current smokers and former smokers from less than 20 years. Life-long time-weighted average of cups/day of coffee and age at diagnosis were broken down in four classes according to the tertiles. DNA adducts were transformed in logarithm, and all values >1 were coded as 1 and otherwise as 0. Genetic biomarkers were coded as 0/1 variables as follows: GSTM1 ("NULL" variant =1, otherwise = 0); GSTP1 ("IA/IA" = 0, otherwise = 1); GSTT1 ("NULL" = 1); NAT1 ("S" = 1); NAT2 ("S" = 1); MPO ("A/A" = 1); COMT ("WW" = 1); MnSOD ("WW" = 1); NQO1 ("MM" = 1); CYP1B1 ("WW" = 0); XRCC1 ("G/G" = 0); XRCC3 ("C/C" = 0); XPD ("A/A" = 0). All the above variables became the predictors in a model of logistic regression in which the dependent variable was 1 for the 15 patients (cases) with AA-related BC (see above) and 0 for the other 184 BC patients (controls). A stepwise selection of independent variables was made using 0.10 as “p-to-enter” and 0.15 as “p-to-remove”. Therefore, from the same sample of 199 cases of BC, two algorithms were obtained (full model and short model) reporting for each regressor the OR with 95% CI and p-value. The best fitting model was chosen with measures of predictive power (R-square and area under the ROC curve) and GOF statistics (Pearson chi-square and Hosmer-Lemeshow test). The criterion was “the higher the better” for the former, and “p-value above 0.05” for the latter. The graphical outputs of the ROC curves were obtained and the AUC was interpreted according to the classification proposed: 0.5 (not informative test); 0.5–0.70 (inaccurate test); 0.7–0.9 (moderately accurate test); 0.9–1.0 (highly accurate test); 1 (perfect diagnostic test) [25]. A statistical test comparing the equality of AUCs was also calculated. Lastly, using the prediction equation we obtained a new variable containing the model-predicted probability of a positive outcome; the same computer program provided the “best cut-off” of predicted probability [26] that maximized the difference between BC patients with or without AA-related disease in both models. Using such value we built the classification table (true positive, false positive, true negative and false negative) from which we calculated sensitivity, specificity, positive and negative predictive values and diagnostic accuracy of each model. Purely statistical measures for comparing two risk prediction models have, however, limited use for medical decision making because they do not incorporate harms and benefits related to treatment decisions arising from the risk prediction model [27]. To evaluate whether clinical use of prediction models, diagnostic tests, and molecular markers would do more good than harm, a simple type of decision analysis (net benefit, NB, approach) has been used [28]. The NB depends on the benefit B, the cost C, the prevalence P of the outcome, and the risk threshold, R, which expresses the model’s classification accuracy, that is the ability of the risk model to assign high risks to cases and low risks to controls. The key quantity is R, which is a function of the harms and benefits of the possible outcomes without detailed specification of harms and benefits [29]. Therefore, the NB to the population of using the risk model is:

$$NBR = (TPR \times P) - \left( \frac{R}{1-R} \right) \times FPR \times (1-P)$$

where:

- **TPR** = true positive rate, also called sensitivity;
- **FPR** = false positive rate, also called one minus specificity;
- **P** = probability of diseases at a given time, also called prevalence;
- **R** = risk threshold, also called model-predicted probability of a positive outcome.

Net benefit can be plotted against a range of R, in what is called a “decision curve”. Decision curves are now widely used in the literature [28]. In the present study, however, wider effects on NB were observed with variations of P rather than of R.
Sample size
In the present one-sample study, the sample size was estimated based on Fisher’s z test assuming a correlation 0.28 (that between DNA adducts and the 0/1 variable “presence/absence of occupational AA-related BC”) and a significance level of 0.05. For a two-sided hypothesis test, the estimated sample size was 98 or 130 patients setting the power at 0.80 or 0.90, respectively [30]. The actual number of BC patients was 199. All statistical analyses were performed with STATA 13.

Results
Table 1 shows the main characteristics of BC patients and their distribution by genotypes (only patients with value set at 1 according to the above definitions). Those with disease related to AAs showed higher level of

| Variables                        | 15 AA-related BC cases | 184 other BC cases |
|----------------------------------|------------------------|-------------------|
| DNA adducts×10⁸ nucleotides (ln) | 1.40 (1.5)             | 0.76 (1.2)        |
| ≥1                               | 11 (73.3)              | 69 (37.5)         |
| Age (years)                      | 60.0 (10.4)            | 63.4 (11.2)       |
| ≤ 56.9 years                     | 7 (46.7)               | 43 (23.4)         |
| 57–65.9 years                    | 3 (20.0)               | 50 (27.2)         |
| 66–70.9 years                    | 3 (20.0)               | 44 (23.9)         |
| ≥ 71 years                       | 2 (13.3)               | 47 (25.5)         |
| Smoking (categories)             |                        |                   |
| Non Smokers & Ex Smokers quitting >20 years | 2 (13.3) | 35 (19.0) |
| Ex Smokers quitting <20 years    | 1 (6.7)                | 60 (32.6)         |
| Current smokers                  | 12 (75.0)              | 87 (47.3)         |
| Pack years (cigarettes smoked lifetime) | 31.5 (12.3) | 35.6 (26.2) |
| ≤ 18.9                           | 1 (6.7)                | 46 (25.1)         |
| 19–32.9                          | 8 (53.3)               | 45 (24.6)         |
| 33–46.9                          | 4 (26.7)               | 46 (24.6)         |
| ≥ 47                             | 2 (13.3)               | 47 (25.7)         |
| Coffee consumption (weighted mean) | 2.1 (1.7)             | 2.4 (2.4)         |
| ≤ 1 cup/day                      | 6 (40.0)               | 71 (38.8)         |
| 2 cups/day                       | 4 (26.7)               | 36 (19.7)         |
| 3 cups/day                       | 3 (20.0)               | 32 (17.5)         |
| ≥ 4 cups/day                     | 2 (13.3)               | 44 (24.0)         |
| Genotypes                        |                        |                   |
| GSTM1                            | 12 (80.0)              | 117 (63.9)        |
| GSTT1                            | 3 (20.0)               | 38 (20.8)         |
| GSTP                             | 5 (33.3)               | 92 (50.3)         |
| NAT1                             | 4 (26.7)               | 60 (32.8)         |
| NAT2                             | 10 (66.7)              | 111 (60.7)        |
| MPO                              | 1 (6.7)                | 6 (3.3)           |
| COMT                             | 7 (46.7)               | 132 (72.1)        |
| MnSOD                            | 4 (26.7)               | 63 (34.4)         |
| CYP1B1                           | 12 (80.0)              | 156 (82.3)        |
| XRCC1                            | 7 (46.7)               | 98 (53.4)         |
| XRCC3                            | 6 (40.0)               | 103 (56.3)        |
| XPD                              | 7 (46.7)               | 113 (61.8)        |

GSTM1 Glutathione S-transferase M1, GSTT1 Glutathione S-transferase T1, GSTP1 Glutathione S-transferase P1, NAT1 N-acetyltransferase isozymes 1, NAT2 N-acetyltransferase isozymes 2, MPO Myeloperoxidase, MnSOD Manganese Superoxide Dismutase, COMT Catechol-O-methyltransferase, CYP1B1 Cytochrome p450 1B1, XRCC1 X-ray repair cross-complementing protein 1, XRCC3 X-ray repair cross-complementing protein 3, XPD xeroderma pigmentosum group D
adducts, lower mean age (with higher percentage of youngest subjects) and higher number of smokers. Except for COMT, GSTM1 and GSTP1 other differences were insignificant.

Table 2 shows the logistic regression analysis obtained with the full and short models (with the subset of predictors chosen by the stepwise selection procedure). As expected from distribution of BC cases, significant ORs were few (DNA adducts and COMT) in both models. Nonetheless, measures of predictive power (R-squares and AUCs) were elevated. The full model seemed more performing, although both models passed the test and were correctly specified.

Table 2 Logistic regression (full and short models)

| Variables | Full model | Short model |
|-----------|------------|-------------|
| Age (years)<sup>a</sup> | | |
| 57–65.9 | 0.12 | 0.99–0.98 | 0.048 |
| 66–70.9 | 0.29 | 0.04–2.25 | 0.239 |
| ≥ 71 | 0.10 | 0.01–1.07 | 0.056 |
| Smoking<sup>b</sup> | | |
| Ex-smokers | 0.05 | 0.00–1.47 | 0.083 |
| Smokers | 1.16 | 0.16–8.60 | 0.887 |
| Coffee consumption<sup>c</sup> | | |
| 2 | 0.63 | 0.09–4.64 | 0.654 |
| 3 | 1.96 | 0.27–14.24 | 0.505 |
| ≥4 | 0.05 | 0.00–0.86 | 0.039 |
| DNA adducts<sup>d</sup> | | |
| ≥1 | 19.20 | 2.52–146. | 0.004 |
| Genotypes<sup>e</sup> | | |
| GSTM1 | 3.01 | 0.51–17.7 | 0.223 |
| GSTT1 | 0.29 | 0.04–1.92 | 0.198 |
| GSTP1 | 0.19 | 0.03–1.08 | 0.061 |
| NAT1 | 0.27 | 0.05–1.52 | 0.139 |
| NAT2 | 2.55 | 0.52–12.6 | 0.25 |
| MPO | 1.38 | 0.03–65.1 | 0.871 |
| COMT | 0.05 | 0.01–0.39 | 0.005 |
| MnSOD | 0.30 | 0.06–1.49 | 0.142 |
| CYP1B1 | 0.27 | 0.03–2.42 | 0.24 |
| XRCC1 | 0.97 | 0.22–4.22 | 0.966 |
| XRCC3 | 0.38 | 0.09–1.68 | 0.203 |
| XPD | 0.31 | 0.05–1.81 | 0.195 |

Reference groups: <sup>a</sup>painents with ≤56.9 years; <sup>b</sup>non smokers and ex-smokers from >20 years; <sup>c</sup>painents with consumption of ≤1 cup/day; <sup>d</sup>painents with <1 (logarithm values) of DNA adducts×10<sup>d</sup> nucleotides; <sup>e</sup>painents with genotype values set at 0 (see text for definition).

Measures of fit for logistic regression

| | Full model | Short model |
|-----------------|------------|-------------|
| R-square | 0.389 | 0.228 |
| Area under the ROC curve | 0.931 | 0.856 |
| Pearson chi-square test: | | |
| p-value | 0.087 | 0.826 |
| no. covariates | 197 | 44 |
| Hosmer-Lemeshow test: | | |
| p-value | 0.130 | 0.042 |
| no. groups | 10 | 10 |

Figure 1 shows the ROC curves obtained for the full and short model; the AUC and its standard error is also reported. A chi-square test comparing the two AUCs gave a value of 4.0438, with <i>p</i> = 0.0443, suggesting that including more variables (algorithm 1) could significantly improve the prediction (e.g. probabilities of occupational AA-related BC). The ROC area of model 1 (= 0.931) fell in the category of highly accurate tests; however, the cost of such high diagnostic accuracy was entering the algorithm 12 genotypes together with DNA adducts and some clinical factors.

Table 3 shows the 15 AA-related BC cases and 184 non-AA-related BC cases, classified as positive and
negative according to the full or short model, always using the “best cut-off” of predicted probability provided by computer program to maximize the difference between BC patients with or without AA-related disease. With respect to the full model, the short one reduced the percentage of cases correctly classified (150/199 = 75% against 182/199 = 92%) by decreasing specificity (137/184 = 75% versus 169/184 = 92%). BC cases classified as ‘positive’ are those to be referred for evaluation by an occupational physician for etiological diagnosis; these patients were 28 (full model) or 60 (short model). Therefore, using 3 instead of 12 genotypes can double the number of patients to be referred for etiologic diagnosis. The other side of the coin was that two out of 15 true AA-related BC patients were classified as ‘false negative’ cases, which like the ‘true negatives’ should not undergo etiologic assessment of their disease. The NB per 100 patients was +4.9 and +4.1 in the left and right panel, respectively, when using panel-specific sensitivity, one minus specificity and the risk threshold R, along with the prevalence of AA-related BC, which was always 0.0754 (= 15/199). Comparing the left and right panel, values of NB were close in spite of divergent values of R.

Table 4 shows the results of a different strategy in which we purposely reduced the “best cut-off” of predicted probability supplied by computer program in order to reduce the false negatives (1 in place of 2). The cost balancing the benefit (decrease of false negatives and increase of true positives) was a higher number of positives (from 28 to 48 with full algorithm; from 60 to 87 with short algorithm) requiring referral to an occupational physician. The NB per 100

| Classified | True | Total | Classified | True | Total |
|------------|------|-------|------------|------|-------|
|            | D    | −D    |            | D    | −D    |
| Positive   | 13   | 15    | 28         | Positive | 13   | 47    | 60 |
| Negative   | 2    | 169   | 171        | Negative | 2    | 137   | 139 |
| Total      | 15   | 184   | 199        | Total   | 15   | 184   | 199 |
| Sensitivity| 86.7%|       | 86.7%      | Sensitivity | 74.5%|
| Specificity| 91.9%|       | 91.9%      | Specificity | 74.5%|
| Positive predictive value | 46.4% | Positive predictive value | 21.7% |
| Negative predictive value | 98.8% | Negative predictive value | 98.6% |
| Correctly classified | 91.5% | Correctly classified | 75.4% |
| Net Benefit per 100 patients | +4.9 | Net Benefit per 100 patients | +4.1 |

The risk threshold R (optimal cut-off point of predicted probability provided by the program) was 0.181 and 0.093 in the left and right panel, respectively. Net Benefit per 100 patients calculated from the above values.
patients was +5.3 and +5.1 in the left and right panel, respectively, when using panel-specific sensitivity, one minus specificity and the risk threshold R, along with the prevalence of AA-related BC, which was always 0.0754 (= 15/199). Comparing the left and right panel, values of NB were close in spite of the divergent values of R.

The little effect on NB with variations of R is also shown in Fig. 2, which depicts NBs as squares and hollow squares (corresponding to R values of 0.181 and 0.093, respectively, see Table 3), or as circles and hollow circles (corresponding to R values of 0.090 and 0.050, respectively, see Table 4). By contrast wider differences and a steep increasing trend of NBs were found with prevalence of AA-related BC going from 0.0 to about 0.14. When prevalence approaches to 0.0, the sign of NBs became negative indicating costs that overwhelm benefits.

Discussion

With the aim of finding a biomarker profile enabling to discriminate AA-related BC from non-AA-related BC, an etiologic prediction model was developed integrating 12 genotypes, DNA adducts, age, smoking and coffee consumption, while using 15 AA-related BC cases. The procedure classified the whole 199 BC patients in 28 positives and 171 negatives. The latter could be left out from etiologic workup, the former are those to be referred for etiological diagnosis. The cases correctly classified were 92% (182/199) and the discriminatory ability was excellent (AUC = 0.93). Examining 3 rather than 12 genotypes the cost of etiologic assessment increased because 60 (instead of 28) positives should receive a further testing (see Table 3, left and right panel). Nevertheless, there were two false negative cases. To overcome this detrimental outcome we used a second strategy (a lower risk threshold) that involved 1 false negative in place of 2 but increased the cost of diagnostic workup since the positives became 48 (instead of 28) or 87 (instead of 60) according to the full or short model, respectively (see Table 4, left and right panel). Nevertheless, there were two false negative cases. To overcome this detrimental outcome we used a second strategy (a lower risk threshold) that involved 1 false negative in place of 2 but increased the cost of diagnostic workup since the positives became 48 (instead of 28) or 87 (instead of 60) according to the full or short model, respectively (see Table 4, left and right panel). With the second strategy, despite the lower percentage of cases correctly classified and regardless of AUC reduction to values (0.7 to 0.9) indicating a moderately accurate test, the benefits were higher than the costs, as it can be seen by comparing the values of NB reported in the Tables 3 and 4.

In our earlier study [20], occupational AAs exposure was found to be positively associated with both BC risk \( (p = 0.041) \) and DNA adducts \( (p = 0.028) \). Since they were not associated with BC risk, DNA adducts were likely biomarkers of exposure. However, the responsible electrophilic substance could not be identified because adducts detected by the nuclease P1 method of 32P–post-labeling are not specific.
As shown in Table 2, many genotypes have given partial regression coefficients (namely, logarithms of ORs) that are not statistically significant. Each could be eliminated without significantly affecting the measures of fit for logistic regression, while the suppression of the whole set had a major effect. In fact, AUC of 0.931 for the full model reduced to 0.69 by entering DNA adducts as single predictor in the logistic regression. This occurs when the variables are strongly related to one another; if one is eliminated the other variables of the group act as substitutes. If, however, the entire group is removed do not remain other variables to compensate the lack of them [31]. In view of the above, we considered all regressors to obtain the predicted probability of a positive outcome, even though it can become difficult to attribute a meaning to each partial regression coefficient.

The etiologic prediction model tool that we have elaborated, enables to divide an initial population into a smaller fraction of “positives” and a larger portion of “negatives”. The former could be referred for further diagnostic assessment, the latter should be no longer considered. The “diagnosis” consists in attributing the disease to an exposure/occupational risk factor; this may happen or might be necessary in several context, such as individual case appraisal, compensation claims, litigation, health authority enquiries. Attributing the disease to an exposure/occupational risk factor may result in several advantages from clinical, epidemiological, individual and public health standpoints. Unfortunately, underreporting to health authorities and under compensation of occupational cancers are well known facts [32–34]. The latter evidence strengthens the need to adopt the second strategy aimed at increase as much as possible the identification of AA-related BC cases. All BC patients are hospitalized at some point in time. Hospital physicians might then face two alternatives when dealing with a tumor, such as BC, with significant incidence and prevalence and with relevant attributable fraction of occupational risk factors: seeking for advice by an occupational health specialist for the patients or manage the case by themselves. A non-selective application of etiologic workup and appraisal would however results in a great loss of clinical and preventive opportunities. In addition to traditional methodology of etiologic diagnosis, a reliable option could therefore be the tool described in this paper that enables discrimination of BC patients with high probability of occupational BC. However, attention should be paid to the underlying risk factors of AA-related bladder cancer. The greater the local degree of industrial development, the higher the chance of occurrence of an occupational disease, and the better the net benefit of using the tool that we have elaborated to ascertain this disease (see Fig. 2).

Validation of model in an external population is an essential next step towards practical use in the clinical setting. External validation requires a multicenter cohort and a prospective collection of data. At the end of the study, the individual characteristics of the validation cohort are multiplied by the regression coefficients of the corresponding variables (those obtained in the internal population) and the products are added to the constant term of the logistic regression. This value quantifies the individual predicted probability of having an AA-related BC. Subsequently, calibration plots are used to graphically explore the association between predicted probabilities and observed proportions: the points should be centered along a 45-degree line in the graph [12].

Conclusions
BC cases with occupational AA-related disease can be individually assessed and stratified based on a predefined molecular biomarker profile. This tool can help ranking BC patients for referrals to an occupational physician for etiologic workup and appraisal. However, practical use in the clinical setting requires validation of the model in another population.

Abbreviations
Aas: Aromatic Amines; BC: Bladder Cancer; COMT: Catechol-O-Methyltransferase; CYP1B1: Cytochrome P450 1B1; FPR1: False positive rate, also called one minus specificity; GSTM1: Glutathione S-Transferase M1 Null; GSTP1: Glutathione S-Transferase P1; GSTT1: Glutathione S-Transferase T1 Null; INAIL: Istituto Nazionale Assicurazione Infortuni Sul Lavoro; ISCO: International Standard Classification Of Occupations; Mnsod: Manganese Superoxide Dismutase; MPO: Myeloperoxidase; NAT1: N-Acetyltransferase 1; NB: Net Benefit; NQO1: NAD(P)H:Quinone Oxidoreductase; P: Probability of diseases at a given time, also called prevalence; Pbs: Peripheral Blood Lymphocytes; R: Risk threshold, also called model-predicted probability of a positive outcome; ROC: Receiver Operator Characteristic; SULT1A1: Sulfotransferase 1A1; T: Time Since Last Exposure; TCE: Time Since First Exposure; TSFE: Time Since First Exposure; TGL: Transgenic Line; TLC: Thin Layer Chromatography; TPR1: True positive rate, also called sensitivity; TSP: Time Since Last Exposure; XPD: Xeroderma Pigmentosum; XPC: Xeroderma Pigmentosum; XRCC1: X-Ray Repair Cross-Complementing Group 1; XRCC3: X-Ray Repair Cross-Complementing Group 3; XTPR: X-Ray Repair Cross-Complementing Group 1; YPC: X-Ray Repair Cross-Complementing Group 3

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Ethics approval and consent to participate
The local ethical committee approved the study (protocol number 2859/9185, 4th September 1996). All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.
Consent for publication
Informed consent was obtained from all individual participants included in the study. All authors have approved the manuscript for submission and consented for publication. Supporting data are available.

Competing interests
The Authors declare that they have no conflict of interest.

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