Pre-treatment risk stratification tools—the end of an era?

In an attempt to predict outcomes and individualize care for prostate cancer (PCa) patients, parallels are drawn between a patient’s current health status and similar cases in the past. To adopt this in clinical practice, risk stratification tools are frequently used. These are, in essence, equations relating multiple factors for a particular individual to the probability or risk for future occurrence of particular outcomes in a certain time period. The terms prognostic and predictive are frequently misunderstood, especially in the field of biomarkers and precision medicine. When a possible interaction (e.g., an intervention) is taken into account, a measurement can be called predictive because it is associated with the impact of a specific therapy. A control group is always needed to evaluate the interaction between treatment benefit and a biomarker or clinical factor (1). Prognostic factors on the other hand provide information on outcomes regardless of any therapeutic intervention (1,2). A biomarker panel can, for example, assign a tumor’s aggressiveness regardless of the treatment that will be provided, where a predictive test could identify which patients will benefit from radiotherapy after biochemical recurrence post-surgery.

The main goal of risk stratification tools is to answer how lethal prostate cancer is, with and without therapy, and how different therapies influence survival outcomes. Most models use similar predictor variables as originally used by D’Amico to differentiate between risk groups: prostate-specific antigen (PSA) level, clinical T-stage, and biopsy Gleason score (3). Adding more variables generally provides more granularity but also increases complexity and impairs usability, as seen in more recent risk scores and nomograms (4-7).

Due to the abundancy of available pre-treatment tools, it can be hard for clinicians and patients to select the appropriate instrument. Zelic et al. recently reported on a head-to-head comparison of nine widely used pretreatment risk stratification tools predicting PCa death and measured their individual performance within the prospectively collected Swedish prostate cancer database (PCBaSe) (8). They found that the Memorial Sloan Kettering Cancer Center (MSKCC) nomogram, Cancer of the Prostate Risk Assessment (CAPRA) score and Cambridge Prognostic Groups (CPG) showed better performance in predicting prostate cancer death than D’Amico risk stratification system and derived tool. However, a great effort is done by the team of Zelic, there are some additional key factors to consider when comparing risk models.

Study population and origin of data

Ideally, the development of risk stratification tools is based on data of a prospective longitudinal cohort study. Information on received diagnostic and staging examinations and treatment modalities should be readily available and indicates generalizability and usefulness of the model. The medical field is continuously evolving and patients recruited a long time ago may have undergone different treatments than recommended by current guidelines. For example, primary androgen deprivation therapy (ADT) monotherapy for high-risk PCa is no longer considered as a contemporary treatment strategy and
nowadays magnetic resonance imaging prior to biopsies is advised. In the Swedish database, the cohort was stratified by treatment and year of diagnosis to tackle this issue when calculating concordance indices (9).

The tools themselves, however, are developed in selected cohorts treated in different time frames. The D’Amico risk groups are based on localized PCa patients who underwent definitive local therapy prior to 1998, thus without accounting for deferred treatment strategies such as active surveillance or watchful waiting (3). This points out the need for exhaustive external validation in different cohorts to confirm the generalizability of the predicted risks.

Predicted outcome

The principal aim is to predict outcomes that are clinically relevant for patients such as survival, morbidity or quality of life measurements. When developing prediction models, ideally the outcome should be assessed while blinded to information about the predictor variables. Otherwise, this knowledge may influence outcome assessment which can lead to a biased estimation of the association between predictor and outcome. This is particularly important for outcomes requiring interpretation, such as PCa specific mortality, radiologic imaging or pathology (5). Pathologists, for example, should be blinded for PI-RADS scores when scoring biopsies to assess the predictive power of pre-biopsy MRI.

Given the high 5-year survival rate for localized PCa, without substantial evidence for superiority of one treatment over another, the CEASAR study group opted to investigate post-treatment functional outcomes based on patient reported outcomes (10). Recently they published a web-based tool to predict functional outcomes after treatment, based on pretreatment characteristics (11,12). Within the online PREDICT prostate tool, findings from this study are also incorporated, however without adapting for pretreatment functional scores (13).

D’Amico risk groups and other derived tools were developed to predict surrogate endpoints such as biochemical recurrence, which do not always have a clear causal relation to relevant patient outcomes (3,6). The CAPRA score predict probability of disease-free survival (which is based on observed disease recurrence rates, i.e., confirmed PSA raise >0.2 ng/mL or additional treatment), only in patients for whom a radical prostatectomy is planned (14). However, even though not developed for this outcome, D’Amico model and the CAPRA score have later been correlated with Cancer Specific Survival (CSS) (15,16).

Crucially, in prediction studies, the effective sample size is determined by the observed outcome events. In the Swedish database, median follow-up time was only 5.8 years, which is relatively short to evaluate PCa specific death (8). A large sample size may be inadequate if only few outcome events are observed.

Predictors

Candidate predictors are typically obtained from patient demographics, medical history, physical examination, disease characteristics and test results (14). Registries contain a large number of potential predictors, but this leads to a higher probability of including weak or useless predictors in the model, overfitting and less user-friendly models (17).

Since the availability and adoption of multi-parametric magnetic resonance imaging (mpMRI) in PCa evaluation, a new generation of risk calculators incorporating mpMRI data have been introduced. Recently, Mortezavi compared conventional, image and biomarker-based risk calculators and concluded that MRI-based predictions outperform conventional models with clinical parameters only in terms of performance (discrimination and calibration) and net benefits (summing up true-positive biopsies and subtracting false-positive biopsies weighted by a factor related to the relative harm of a missed cancer versus an unnecessary biopsy) to distinguish clinical significant PCa (9).

Moreover, prognostic models may focus on a cohort of patients who are yet to receive any kind of treatment and where treatment can be included as a potential prognostic factor, as seen with the PREDICT prostate model (5). However, caution is needed when including treatment as a prognostic factor because administration of treatment should be standardized to prevent bias of confounding by indication and the predictive impact of treatment is generally small compared to other predictive variables (14).

In Table 1 a subset of the criteria for Transparent reporting of a multivariable prediction model for individual prognosis or diagnosis (TRIPOD) (17) is used to summarize key characteristics of the three best performing tools according to Zelic et al., compared with the more recent PREDICT prostate and D’Amico risk classification. Each of these tools use at least the same predictor variables as described by D’Amico, but additionally there are some differences. All, except for D’Amico differentiate between primary and secondary Gleason grade thus discriminating...
between 3+4 versus 4+3. CAPRA, MSKCC and PREDICT prostate also account for percentage of positive biopsies as surrogate for tumor volume. This variable is optional for the latter two models, but it is necessary to calculate CAPRA scores. Therefore, CAPRA scores can only be calculated when 6 or more biopsies have been obtained, which was contemporary practice before the introduction of targeted biopsies (5,6). Consequently, caution is advised when calculating CAPRA scores with modern biopsy techniques.

PREDICT prostate is the only model that includes comorbidities as a predictive variable, but the predictive effect was only seen in relation to non-prostate cancer specific death (18). A model that incorporates a holistic view of the patients and predicts OS instead of CSS is obviously of greater interest for patients and clinicians.

### Table 1: Comparison of the three best performing prognostic tools according to Zelic, compared with D’Amico and predict prostate

<table>
<thead>
<tr>
<th>System</th>
<th>Study population</th>
<th>Outcome</th>
<th>Predictors</th>
<th>Statistical analysis methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>D’Amico (3)</td>
<td>1,872 men treated between January 1989 and October 1997 by an RP (n=888) or interstitial radiation with or without neoadjuvant ADT (n=218), or RT (n=766)</td>
<td>PSA failure (at least 3 consecutive rising PSA values, obtained at least 3 months apart)</td>
<td>PSA level, biopsy Gleason score, T-stage (AJCC 1992)</td>
<td>Cox regression</td>
</tr>
<tr>
<td>CAPRA (6)</td>
<td>1,439 RP patients (from CaPSURE™ database - Cancer of the Prostate Strategic Urologic Research Endeavor) diagnosed between 1992 and 2001 with T1c-3a, N0/x, M0/x, without (neo)adjuvant RT of ADT. Only with known PSA, Gs, cT-stage and at least sextant biopsy, at least two follow-up PSAs or evidence of additional treatment &gt;6 months after RP</td>
<td>Disease-free survival (two consecutive PSA values ≥0.2 ng/mL postoperatively or additional treatment &gt;6 months after RP)</td>
<td>PSA, Gleason score, T stage, PPB, age at diagnosis</td>
<td>Cox proportional hazard</td>
</tr>
<tr>
<td>CPG (4)</td>
<td>10,139 patients diagnosed with ICD-10 code C61 between 1 January 2000 and 31 December 2010. Only cases with complete staging, grades and PSA; follow-up and survival data. (789 prostate cancer deaths and 2,610 overall deaths) (censoring September 2013)</td>
<td>Prostate-cancer-specific mortality (PCSM)</td>
<td>Age, PSA, ISUP score, T stage</td>
<td>Cox proportional hazard; Fine-Gray test was applied for competing-risk regressions</td>
</tr>
<tr>
<td>PREDICT prostate (5)</td>
<td>10,089 patients diagnosed between 1 January 2000 and 31 December 2010. Only cases with complete staging, grades and PSA; follow-up and survival data. Comorbidity scores are also included (censoring September 2016)</td>
<td>PCSM and non-PCa mortality (NPCM)</td>
<td>Age, PSA, histological grade group, clinical T-stage, primary treatment type, comorbidity (only for NPCM), PPB (optional)</td>
<td>Separate multivariable cox proportional hazard for PCSM and NPCM</td>
</tr>
<tr>
<td>MSKCCC (7)</td>
<td>10,000 PCa patients treated at MSK</td>
<td>PCSM</td>
<td>Age, PSA, histological grade group, clinical T-stage, PPB (optional)</td>
<td>N/A</td>
</tr>
</tbody>
</table>

RP, radical prostatectomy; ADT, androgen deprivation therapy; RT, radiation therapy; PSA, prostate specific antigen; Gs, Gleason score.

### Model specification

Deciding the appropriate treatment modality is often influenced by survival predictions. Guidelines recommend radical prostatectomy only in patients with few comorbidities and a >10 years life expectancy (19). This implies low PCa specific mortality and consequently, other causes of death may occur prior to death of prostate cancer (e.g., death of heart failure can precede prostate cancer specific mortality). These competing risk events can influence the supposed benefit of active treatment and consequently influence the decision process (20). A standard Cox proportional hazard model leads to an increased risk prediction compared to a Fine and Gray model due to the lack of accounting for competing risks, which is especially important in an elderly population (21). Thus, it is important to acknowledge the effect of a standard Cox proportional hazard model on survival predictions.
model developed within a radical prostatectomy population, in which patients de facto have a better health status, when generalizing it to a newly diagnosed PCa population with their typical competing risks.

**Machine learning (ML) predictions requires a paradigm shift**

When using prediction tools, clinicians often fill out all the necessary variables in an online calculator or paper based nomogram or they select the corresponding risk group. Crucial to widespread adoption is the usability. Overall, compact models with few variables are more frequently used (17). Besides usability, utility plays an important role and can be achieved when predictions can be adequately interpreted and translated into clinical practice.

A multivariable approach also enables researchers to investigate and quantify whether new (bio)markers or investigations have an added predictive value over traditionally used clinic-pathological variables (14). Notice that a large dataset is necessary to provide evidence for significant added value of a test. But if this new test is not incorporated in prognostic models, its adoption might be hampered because the perception might live that there is not enough predictive value in it and consequently no evidence of significant added predictive value can be generated.

A prognostic model might be used to provide insight into causality of the predictor variable and the studied outcome, however this is not equivalent to etiological research. The latter has a goal to explain, whether an outcome can be attributed to a risk factor, with adjustment for confounding factors (14). In a prognostic study, all variables which are potentially associated with the outcome, not necessarily causally, can be considered. All causal factors are predictors but not every predictor is a cause (14). PSA, for example is an example of a predictive but non-causal factor for disease recurrence. This is of importance since both prognostic and etiological research are based on similar design and analysis. Conversely, etiological research, uses statistical inference to characterize the relations between the data and the outcome variable. Often, these two go hand-in-hand since causality and the understanding of underlying drivers of disease are of specific interest to develop new therapeutic strategies. Figure 1 shows the relationship between inferential and predictive statistics.

In a visionary commentary by Vickers the question is raised “why can’t nomograms be more like Netflix?” (22). They describe four major advantages of the online streaming platform Netflix-like algorithms compared to traditional nomograms and make the bold statement that medical prediction can be done using technologically superior methods.

Traditional risk prediction tools are based on regression analyses of manually acquired data. ML on the other hand is a form of artificial intelligence (AI) and refers to computer systems that learn from raw data with some degree of autonomy (23). The weaknesses of traditional approaches may be overcome by directly incorporating electronic health record (EHR) data into ML models to augment decision making through different phases (23).

Multiple challenges, some similar as seen in traditional prediction model development, are blocking today’s adoption of ML techniques. Data standardization, model interpretability, implementation and monitoring as well as ethical issues must be carefully considered (23).

A real paradigm shift would happen if we move away from predictions based on relatively small data obtained by “manual” observations to large amounts of data produced by high-throughput instruments analyzed by ML “black box” models that might produce more accurate predictions. ML models are often perceived as black boxes since the algorithm implicitly learned the features, possibly unknown for physicians, that were most predictive based on the data (24).

Regardless of how the model is built, it should be externally validated an the clinical effectiveness must be assessed (25). The TRIPOD steering group together with a large group of experts is developing guidelines for prediction models that are based on AI or ML which will contribute to improvements in the design and reporting of such studies (26).

**Conclusions**

A holistic view of PCa patients together with an integrated approach of the tumor combining advanced diagnostics and treatment strategies will enable more accurate and relevant predictions. Newer models, such as PREDICT prostate, incorporate already comorbidities and calculate the effect of radical versus conservative treatment, and enable the adoption of additional factors if proven prognostic. However, it is arguable that complex concepts as comorbidities can hardly be summarized in one binary variable. Further, the settings in which therapy will be delivered cannot be underestimated. Surgeon’s experience, hospital volume and a dedicated team are already shown...
to have an impact on surgical patient’s outcomes (27). It becomes clear that the abundance of data generated combined with ongoing insights and novel technologies calls for a change in the methodology of predicting outcomes that matters for PCa patients, with overall survival as a top priority.

As long as ML models are still under development, the uro-oncological community may be helped by analyses as done by Zelic et al. to choose a suitable model for predicting PCa patient’s survival.

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