askMUSIC: Leveraging a Clinical Registry to Develop a New Machine Learning Model to Inform Patients of Prostate Cancer Treatments Chosen by Similar Men

Gregory B. Auffenberg a, Khurshid R. Ghani b, Shreyas Ramani c, Etiwo Usoro c, Brian Denton b,d, Craig Rogers e, Benjamin Stockton f, David C. Miller b, Karandeep Singh c,g,h,*

for the Michigan Urological Surgery Improvement Collaborative

Abstract

Background: Clinical registries provide physicians with a means for making data-driven decisions but few opportunities exist for patients to interact with registry data to help make decisions.

Objective: We sought to develop a web-based system that uses a prostate cancer (CaP) registry to provide newly diagnosed men with a platform to view predicted treatment decisions based on patients with similar characteristics.

Design, setting, and participants: The Michigan Urological Surgery Improvement Collaborative (MUSIC) is a quality improvement consortium of urology practices that maintains a prospective registry of men with CaP. We used registry data from 45 MUSIC urology practices from 2015 to 2017 to develop and validate a random forest machine learning model. After fitting the random forest model to a derivation cohort consisting of a random two-thirds sample of patients after stratifying by practice location, we evaluated the model performance in a validation cohort consisting of the remaining one-third of patients using a multiclass area under the curve (AUC) measure and calibration plots.

Results and limitations: We identified 7543 men diagnosed with CaP, of whom 45% underwent radical prostatectomy, 30% surveillance, 17% radiation therapy, 5.6% androgen deprivation, and 1.8% watchful waiting. The personalized prediction for patients in the validation cohort was highly accurate (AUC 0.81).

Conclusions: Using clinical registry data and machine learning methods, we created a web-based platform for patients that generates accurate predictions for most CaP treatments.

Patient summary: We have developed and tested a tool to help men newly diagnosed with prostate cancer to view predicted treatment decisions based on similar patients from our registry. We have made this tool available online for patients to use.

© 2018 European Association of Urology. Published by Elsevier B.V. All rights reserved.

* Corresponding author. Department of Learning Health Sciences, University of Michigan Medical School, Ann Arbor, MI 48109, USA. Tel. +1 734 9361649; Fax: +1 734 6473914. E-mail address: kdpsingh@umich.edu (K. Singh).
1. Introduction

Physicians and other health care stakeholders are increasingly using clinical data registries to enhance their own decision-making and improve health care quality [1-3]. Recognizing their potential, many national medical organizations have invested heavily in clinical registries as a foundation for quality improvement efforts such as the American College of Surgery’s National Surgical Quality Improvement Program and the American Urological Association’s AQUA registry. The Centers for Medicare and Medicaid Services established the qualified clinical data registries program as part of the Physician Quality Reporting System in 2014 and further elevated registries in the 2015 Medicare Access and CHIP Reauthorization Act to provide a greater financial incentive for participation in registries, recognizing that they can serve as a unique tool “to foster improvement in the quality of care provided to patients” [4].

Although registry initiatives have gained prominence during a period notable for increasing focus on patient access to personal health information [5], there are few examples whereby patients can directly interact with clinical registries. The information in these large data repositories, if presented appropriately, may represent an important avenue for patients to find individualized, evidence-based answers to questions about a new diagnosis or prescribed treatment. Strategies commonly used to distill data into provider-facing tools to guide decision-making (eg, predictive modeling, machine learning) [1,6-8] may prove powerful for generating patient-facing content from registry data that can specifically address patient-centered questions. Men with localized prostate cancer often face multiple viable treatment options. In the setting of uncertainty, high-quality decisions are those that are informed, values-based, and implemented without undue delay [9].

In this context, we used data from a large clinical registry of men diagnosed with prostate cancer maintained by the Michigan Urological Surgery Improvement Collaborative (MUSIC) and input from patient advocates to develop and validate a model to help patients focus their self-education on the treatments most likely to be recommended by their urologist. Although use of such a model will not replace the shared decision-making process, it may complement the information that patients receive from traditional educational materials, potentially reducing decisional uncertainty for patients. As a second aim, we worked to integrate this model into a web-based platform to generate a user-friendly interface that patients can use to interact with the predictive model to augment their decision-making.

2. Patients and methods

2.1. Data source

MUSIC was established in 2011 in partnership with Blue Cross Blue Shield of Michigan. The quality improvement collaborative currently comprises 45 diverse community and academic urology practices representing approximately 90% of the urologists in the state. For all men seen in MUSIC practices who undergo a prostate biopsy, trained data abstractors prospectively enter a standardized set of data elements into the registry database. These elements include demographic and clinicopathologic information related to prostate biopsies, subsequent diagnoses, treatments, and follow-up care, from which both the patient characteristics and treatment strategies are derived. Prior reports have described the data acquisition and quality control activities for MUSIC, which include annual data audits at each practice and validation analyses based on insurance claims [10,11]. Each MUSIC practice obtained an exemption or approval for collaborative participation from a local institutional review board.

2.2. Study cohort

The cohort in the present study included all men in the MUSIC registry newly diagnosed with prostate cancer between November 4, 2015 and November 3, 2017 for whom a subsequent treatment decision was documented. Although MUSIC registry data for prostate cancer date back to 2011, we excluded data from before 2015 because increasing utilization of active surveillance among patients with favorable-risk prostate cancer may lead to outdated treatment probabilities if older data are included.

We excluded patients with metastatic disease using clinical staging information documented within 150 d of the initial biopsy and patients with a missing date of birth or Gleason score or with a weight outside the range of 80–700 lbs. We also excluded patients who had undergone rare treatments, including chemotherapy, cryotherapy, high-intensity focused ultrasound, immunotherapy, and radical cystoprostatectomy.

We divided patients into derivation and validation cohorts using 2:1 random sampling of the overall cohort, stratified by practice location. Thus, each of the 45 urology practices was represented in both the derivation and validation cohorts.

2.3. Model development

Using age at biopsy, prediagnosis prostate-specific antigen level, number of positive cores on biopsy, total cores taken at biopsy, and weight as continuous predictors; primary, secondary, and overall biopsy Gleason scores as numerical values; and history of myocardial infarction and diabetes status as binary predictors, we fitted a multinomial random forest model to predict the probability of receiving a given primary treatment. Random forest is a nonparametric method described by Leo Breiman for classification (binary or multinomial outcome) and regression (continuous outcome) problems [12].

The predicted primary treatment outcomes included: radical prostatectomy, radiation therapy (either external beam, brachytherapy, or both), primary androgen deprivation therapy, active surveillance, and watchful waiting. Active surveillance and watchful waiting are coded as distinct separate entities in the registry, as described elsewhere [13]. Additional details are discussed in the Supplementary material.

2.4. Model validation

After fitting the random forest model using the derivation cohort, the discrimination of the model was evaluated in the validation cohort using a multiclass area under the curve (AUC) measure [14]. Model calibration was evaluated using a calibration plot comparing deciles of predicted probabilities for each outcome with observed probabilities.

2.5. Missing data

Variables with missing values were assumed to carry information. During model training, optimal binary splits were determined by minimizing the error using nonmissing data. After a variable split was
determined, missing values for that variable were assigned to the direction minimizing the error. When generating predictions, missing values followed the assigned direction [15].

2.6. Development of the askMUSIC web platform

The askMUSIC platform was developed using the Shiny web app platform in the R language. The website was designed with input from the study team and four patients with prostate cancer. Patients reviewed the initial version of the tool and communicated their opinions and recommendations first via a conference call and then by e-mail. This was facilitated by a urologist (G.B.A.) and an informatician (K.S.).

2.7. Software

We used R 3.4.3 for all analyses [16]. The random forest model was implemented using the h2o R package [17], which links to the h2o Java program (version 3.16.0.2).

3. Results

We identified 7543 men with newly diagnosed localized prostate cancer who met the inclusion and exclusion criteria (Fig. 1). Of these men, 3413 (45%) underwent radical prostatectomy, 2289 (30%) active surveillance, 1280 (17%) radiation, 422 (5.6%) androgen deprivation therapy, and 139 (1.8%) watchful waiting. After stratifying by practice location, we randomly assigned 5016 men to the derivation cohort and 2527 to the validation cohort. There was a statistically significant but clinically minor difference in biopsy Gleason score between the two cohorts (Table 1). The difference in treatment choices between the two cohorts was not statistically significant (Table 2).

3.1. Model discrimination and calibration

The model achieved excellent discrimination, with an overall multiclass AUC of 0.81 in the validation cohort (Table 3). The easiest treatments to distinguish from one another were active surveillance and androgen deprivation therapy, while the most difficult to distinguish were active surveillance and watchful waiting. The model was well calibrated in the validation cohort (Fig. 2).

3.2. Model characteristics

Although the model parameters are not readily shareable in a random forest, the relative importance of variables can be derived using a permutation method [18]. Age is the most important variable, followed by the number of positive cores and overall biopsy Gleason score \( p < 0.001 \); Supplementary Table 1). Relationships between the predictors and treatment options are demonstrated using a partial dependence plot in Supplementary Figure 1. In general, higher age is associated with a lower probability of radical prostatectomy and a higher probability of alternative strategies. A higher number of positive cores and a lower number of total cores are both associated with greater probability of undergoing radical prostatectomy. Higher biopsy Gleason score is associated with a lower probability of active surveillance (Supplementary Figs. 1 and 2).

3.3. Model deployment using the askMUSIC platform

We held a teleconference with four patient advocates who are part of the MUSIC collaborative on August 5, 2016 to obtain feedback on the presentation of the model results. The key suggestions that have been incorporated into the tool are summarized in Supplementary Table 2. The model is accessible to patients at http://ask.musicurology.com (Fig. 3). A link to its code is available in Supplementary Table 3.

4. Discussion

We developed a multinomial random forest model to predict the probability of receiving a given treatment for localized prostate cancer based on common clinicopathologic and
Table 1 – Baseline characteristics for the model derivation and validation cohorts

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Derivation cohort</th>
<th>Validation cohort</th>
<th>p value *</th>
<th>Missing (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients (n)</td>
<td>5016</td>
<td>2527</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median age, yr (IQR)</td>
<td>66 (60–71)</td>
<td>66 (60–71)</td>
<td>0.787</td>
<td>0.0</td>
</tr>
<tr>
<td>Bopsy Gleason score, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤6</td>
<td>1762 (35)</td>
<td>874 (35)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 + 4</td>
<td>1639 (33)</td>
<td>790 (31)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 + 3</td>
<td>733 (15)</td>
<td>428 (17)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>502 (10)</td>
<td>239 (9.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>336 (6.7)</td>
<td>183 (7.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>44 (0.9)</td>
<td>13 (0.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median PSA, ng/ml (IQR)</td>
<td>6.1 (4.6–8.9)</td>
<td>6.2 (4.7–9.1)</td>
<td>0.198</td>
<td>3.2</td>
</tr>
<tr>
<td>Median positive cores, n (IQR)</td>
<td>4 (2–6)</td>
<td>4 (2–6)</td>
<td>0.825</td>
<td>1.6</td>
</tr>
<tr>
<td>Median total cores, n (IQR)</td>
<td>12 (12–12)</td>
<td>12 (12–12)</td>
<td>0.550</td>
<td>1.6</td>
</tr>
<tr>
<td>Diabetes mellitus, n (%)</td>
<td>648 (13)</td>
<td>341 (14)</td>
<td>0.507</td>
<td>0.0</td>
</tr>
<tr>
<td>History of MI, n (%)</td>
<td>161 (3.2)</td>
<td>87 (3.4)</td>
<td>0.640</td>
<td>0.0</td>
</tr>
<tr>
<td>Median weight, lbs (IQR)</td>
<td>200 (177–225)</td>
<td>200 (177–225)</td>
<td>0.997</td>
<td>0.0</td>
</tr>
</tbody>
</table>

IQR = interquartile range; PSA = prostate-specific antigen; MI = myocardial infarction.
* A χ² test was used to compare categorical variables and a Wilcoxon test for continuous variables.

Table 2 – Primary treatment received by cohort

<table>
<thead>
<tr>
<th>Primary treatment</th>
<th>Derivation cohort Patients, n (%)</th>
<th>Validation cohort Patients, n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active surveillance</td>
<td>1535 (31)</td>
<td>754 (30)</td>
</tr>
<tr>
<td>Androgen deprivation therapy</td>
<td>283 (5.6)</td>
<td>139 (5.5)</td>
</tr>
<tr>
<td>Radiation</td>
<td>829 (17)</td>
<td>451 (18)</td>
</tr>
<tr>
<td>Radical prostatectomy</td>
<td>2263 (45)</td>
<td>1150 (46)</td>
</tr>
<tr>
<td>Watchful waiting</td>
<td>106 (2.1)</td>
<td>33 (1.3)</td>
</tr>
</tbody>
</table>

* p = 0.088, χ² test.

Table 3 – Model discrimination between differing primary treatments

<table>
<thead>
<tr>
<th>Primary treatments</th>
<th>Validation cohort AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active surveillance vs androgen deprivation therapy</td>
<td>0.96</td>
</tr>
<tr>
<td>Active surveillance vs radiation</td>
<td>0.89</td>
</tr>
<tr>
<td>Active surveillance vs radical prostatectomy</td>
<td>0.90</td>
</tr>
<tr>
<td>Active surveillance vs watchful waiting</td>
<td>0.64</td>
</tr>
<tr>
<td>Androgen deprivation therapy vs radiation</td>
<td>0.67</td>
</tr>
<tr>
<td>Androgen deprivation therapy vs radical prostatectomy</td>
<td>0.84</td>
</tr>
<tr>
<td>Androgen deprivation therapy vs watchful waiting</td>
<td>0.80</td>
</tr>
<tr>
<td>Radiation vs radical prostatectomy</td>
<td>0.68</td>
</tr>
<tr>
<td>Radiation vs watchful waiting</td>
<td>0.80</td>
</tr>
<tr>
<td>Radical prostatectomy vs watchful waiting</td>
<td>0.92</td>
</tr>
<tr>
<td>Multiclass AUC</td>
<td>0.81</td>
</tr>
</tbody>
</table>

AUC = area under the curve.

demographic factors. The model exhibited excellent discrimination and calibration between treatment options in the validation cohort. Integration of the model into a web-based platform allows patients to access this tool on the Internet.

Predictive modeling has long been leveraged to aid physicians looking to use large data sets to augment real-time decision-making [1,2,6,8]: applying these principles to create patient-facing models may represent a meaningful way to provide patients with insights from registry data while also addressing data security issues. Merging high-quality registry data, random forest predictive modeling, and a web platform to allow simple interaction with the models may help patients to rapidly identify the information in a registry most relevant to their situation to augment decision-making. Furthermore, as patients using this tool interact with an interface to query a model as opposed to raw data, it should largely eliminate many of the usability and data security concerns that may arise from allowing interactions with raw data. This tool may provide patients with an objective source of additional information to inform decision-making. Examples of possible uses are as follows: (1) men could use the model to gain perspective on the treatment options that their physician is likely to recommend; (2) the model may enable patients to ask physicians about discrepancies in individual recommendations in comparison to treatments received by similar men as they work to make a decision about what is the best road forward for management; and (3) physicians wanting a patient to seriously consider an alternative management strategy to the patient’s initial choice may benefit from referring patients to the model so they can understand how common certain strategies are for a given stage of disease.

Our findings should be considered in the context of several limitations. First, predictive models learned from human behavior can reinforce human biases [19]. For instance, if active surveillance is appropriate but underutilized in the data, then a model derived from the data may encourage continued underutilization of active surveillance. Similarly, primary androgen deprivation therapy monotherapy is generally not an appropriate treatment for localized prostate cancer, so the model finding that similar men receive this treatment does not necessarily mean that it is medically appropriate. We have tried to avoid treatment suggestions that are based on out-of-date or nonrepresentative practice patterns by discarding older data before model fitting and including all MUSIC practices in the derivation cohort. Nonetheless, the model does not account for personal preferences with respect to survival and the side effects of different treatments, so while this tool has potential to serve as an important support to decision-making, it should not replace physician-led counseling and
Fig. 2 – Calibration plot for each treatment in the validation cohort, with shaded 95% confidence intervals.

Fig. 3 – Screenshot of the askMUSIC web platform.
shared decision-making. Second, how patient use of the predictive model impacts their decision-making is not yet known. In the absence of an objectively correct treatment decision, there is reason to believe that aggregated patient data—either directly or through a model—may be useful in supporting decision-making [20]. In the face of an overwhelming amount of information available online, the model may help patients to focus their self-education on a narrower set of treatment options. Third, we have not externally validated this model outside the state of Michigan. Both our derivation and validation cohorts are representative of overall contemporary practice patterns within MUSIC. Finally, the model performed poorly in discriminating between active surveillance and watchful waiting. Although these management options have different definitions [21], these terms are sometimes applied interchangeably and may explain some of the difficulty with discrimination.

Despite these limitations, our work has important implications. Treatment recommendations can vary substantially across different risk strata and there is often more than one viable treatment option [22]. The last 10–20 yr have been notable for significant changes in how often various management strategies are used for prostate cancer [23]. Deciding on a course of treatment may lead to significant anxiety for patients [24] and may in part explain why many patients favor overtreatment [25,26] in the face of pressure from family members and physicians to select more aggressive treatments than may be warranted [27]. One approach that has proven successful in leveraging patient experiences to drive a better understanding of treatment options and decision-making about health is the use of patient-led online communities [28]. Providing patients with a tool to help them understand decisions made by similar men serves a similar purpose, although the effect of the tool on patient decision-making remains to be seen.

In the future we plan to disseminate the askMUSIC tool across our collaboration. Understanding how the tool impacts patient decision-making and decision uncertainty will be an important next step, and particularly whether the model leads patients to choose less aggressive treatments for low-risk cancers or vice versa.

5. Conclusions

We have developed a predictive model using the random forest machine learning method that uses clinicopathologic and demographic characteristics to provide an individualized prediction of treatment in the state of Michigan. Predictive modeling may represent an important mechanism for unlocking information in data registries to allow patients to learn directly from these valuable repositories.

The abstract on which this manuscript is based was presented at the American Urological Association 2017 annual meeting.

Author contributions: Karandeep Singh had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Study concept and design: Auffenberg, Ghani, Usoro, Denton, Rogers, Stockton, Miller, Singh.

Acquisition of data: Auffenberg, Singh.

Analysis and interpretation of data: Auffenberg, Ghan, Usoro, Ramani, Denton, Rogers, Stockton, Miller, Singh.

Drafting of the manuscript: Auffenberg, Singh.

Critical revision of the manuscript for important intellectual content: Auffenberg, Ghan, Usoro, Ramani, Denton, Rogers, Stockton, Miller, Singh.

Statistical analysis: Ramani, Singh.

Obtaining funding: Auffenberg, Denton, Miller, Singh.

Administrative, technical, or material support: Usoro.

Supervision: Singh, Ghan.

Other: None.

Financial disclosures: Karandeep Singh certifies that all conflicts of interest, including specific financial interests and relationships and affiliations relevant to the subject matter or materials discussed in the manuscript (e.g., employment/affiliation, grants or funding, consultancies, honoraria, stock ownership or options, expert testimony, royalties, or patents filed, received, or pending), are the following: Gregory B. Auffenberg has received funding from the National Cancer Institute (grant 1T32-CA180984). Khurshid R. Ghan has received contract support from Blue Cross Blue Shield of Michigan for serving as the co-director of the Michigan Urological Surgery Improvement Collaborative and serving as a consultant for Boston Scientific Corporation and Lumenis. Shreyas Ramani and Etioowo Usoro have received salary support from an MCubed grant from the University of Michigan. David C. Miller has received contract support from Blue Cross Blue Shield of Michigan for serving as the director of the Michigan Urological Surgery Improvement Collaborative. Karandeep Singh has received grant support from the National Institute of Diabetes and Digestive and Kidney Diseases (grant 5K12DK111011). The remaining authors have nothing to disclose.

Funding/Support and role of the sponsor: This work was supported by Blue Cross Blue Shield of Michigan. The sponsor played no direct role in the study.

Acknowledgments: The corresponding author would like to thank all the support staff from the Michigan Urological Surgery Improvement Collaborative.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j.eururo.2018.09.050.

References


