Budget Impact Model: Epigenetic Assay Can Help Avoid Unnecessary Repeated Prostate Biopsies and Reduce Healthcare Spending

Wade Aubry, MD; Robert Lieberthal, PhD; Arnold Willis, MD; Grant Bagley, MD, JD; Simon M. Willis III, MS; Andrew Layton, BA

Background: The diagnosis of prostate cancer involves invasive, sometimes harmful, procedures that can entail negative quality-of-life implications to individuals and high additional costs to the US healthcare system when these procedures result in retesting and iatrogenic harms. It is estimated that $1.86 billion is spent annually on prostate-specific antigen (PSA) testing alone. An advanced epigenetic molecular diagnostic test that uses methylation-specific polymerase chain reaction to assess the DNA methylation status of GSTP1, APC, and RASSF1 genes associated with oncogenesis enables a higher degree of accuracy (previously unattainable through prostate biopsy procedures alone) and produces clinical, financial, and health benefits by reducing the number of medically unnecessary and costly repeated biopsies that are part of today’s standard of care.

Objectives: The purpose of this study is to quantify, using a budget impact model, the effect of a relatively new epigenetic assay on healthcare costs for commercial health plans that reimburse for the assay, by avoiding unnecessary repeated prostate biopsy procedures.

Methods: A budget impact model was developed to test the hypothesis that the epigenetic assay can produce cost-saving benefits to health plans, as well as clinical benefits to urologists and patients with prostate cancer, by providing guidance on how to offer patients more appropriate, and less costly, care. The budget impact model is presented from the perspective of a hypothetical commercial health plan, and direct costs are calculated over a 1-year time horizon, using 2013 Medicare fee-for-service rates. Using a plan of 1 million members, the model compares 1-year costs in a “reference scenario,” in which the epigenetic assay is not used for the screening and diagnosis of prostate cancer, to costs in a “new scenario,” in which the epigenetic assay is used to distinguish true-negative prostate biopsy results from false-negative biopsy results.

Results: Based on this analysis, administering the epigenetic assay to patients with histopathologically negative biopsies would result in a reduction of 1106 unnecessary biopsies for a health plan with 1 million members. The total 1-year cost of repeated prostate cancer biopsies to the health plan was found to be $2,864,142 in the reference scenario and $2,333,341 in the new scenario. This translates to a total budget impact, or an annual savings, of $530,801 to the plan. The total diagnostic cost was calculated to be $2584 per patient in the new scenario (using the genetic assay) compared with $3172 per patient in the reference scenario (that did not use the assay), resulting in a savings of $588 per patient management.

Conclusion: This analysis shows that the net cost to a commercial health plan with 1 million members would be reduced by approximately $500,000 if patients with histopathologically negative biopsies were managed with the use of the epigenetic assay to differentiate patients who should undergo repeated biopsy and those who should not. Using this genetic-based assay can save costs to health plans and to the US healthcare and improve the clinical management of patients with elevated PSA levels.

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Prostate cancer is the most frequently detected cancer in men, and 1 of 6 men will be diagnosed with prostate cancer during their lifetime based on Medicare enrollment data. In the United States, approximately 19 million men annually are screened by prostate-specific antigen (PSA) testing, resulting in approximately 4.7 million abnormal PSA test results (≥4.0 ng/mL) leading to approximately 1.3 million biopsy procedures. According to the National Cancer Institute, 241,740 men are diagnosed with prostate cancer annually, and 28,170 prostate cancer–related deaths were reported in 2012. Although some forms of prostate cancer are deadly, many forms are low grade and can be managed by active surveillance. Aggressive variants of prostate cancer can be one of the deadliest cancers in men, and accurate diagnosis and follow-up remain a challenge and come at a considerable cost to the US healthcare system.

Clinical Burden

Despite the recent controversy that was raised by the US Preventive Services Task Force (USPSTF) findings on PSA testing, leading to their recommendations to stop routine PSA-based screening, the American Urological Association (AUA) continues to recommend the PSA blood test, along with digital rectal examination (DRE), for screening men at risk for prostate cancer. Screening has led to a shift of detecting earlier-stage disease, resulting in an increased likelihood for curative treatment. If screening is eliminated, urologists fear an increased incidence of advanced cancers and an increase in healthcare costs to effectively treat these patients. Today, urologists typically perform a biopsy for high-risk patients with a rising PSA and for patients with a PSA level ≥4.0 ng/mL, obtaining approximately 10 to 12 needle-core tissue samples according to the current standard of care.

Of note, an abnormal PSA result can often be caused by factors other than cancer, including infection, inflammation, or other benign conditions, such as benign prostatic hyperplasia. This leads to the inclusion of many men with no cancer among those who are being subjected to prostate biopsies (ie, false-positive PSA screening). The rate of cancer detection in men undergoing prostate biopsies is approximately 30%, and approximately 75% of men who have undergone biopsies have negative prostate biopsy results.

An elevated PSA and/or abnormal DRE identify men at high risk for prostate cancer, and, as a result, many of these men will undergo a biopsy procedure. However, because of the nature of random and limited sampling of the prostate, many cancers are undetected by histopathologic review. Studies by urology and pathology opinion leaders report that initial prostate biopsy histopathology has a 20% to 30% false-negative rate.

Molecular Testing

With such high costs to the US healthcare system, as well as negative quality-of-life implications to patients,
A hypothetical plan using patient age-groups between ages 40 and 64 years (similar to those in commercial health plans). An additional sensitivity analysis was conducted for a hypothetical plan consisting of patients aged ≥65 years (similar to a Medicare health plan), using methods identical to the base case, with the exception of the patient ages and corresponding PSA rates. Biopsies and costs for patients younger than age 40 years or older than age 74 years were not included in the analysis.

The model's base-case analysis was conducted for a hypothetical plan consisting of patients aged ≥65 years (similar to a Medicare health plan). Biopsies and costs for patients younger than age 40 years or older than age 74 years were not included in the analysis.

The cost analysis was based on total costs of 1 year in the course of prostate cancer screening and evaluation through prostate biopsy. Costs to the health plan are assumed to be equal to the Medicare FFS rates, which provide a conservative benchmark for reimbursement rates paid for by other health plans.

The model allows for the simulation of the current (reference scenario) and a counterfactual reality (new scenario). In the reference scenario, the model uses current clinical patterns of care to simulate the treatment of men at risk of prostate cancer in the reference scenario; a molecular assay was not utilized for prostate cancer detection. In the new scenario, men at risk for repeated biopsy are evaluated with the epigenetic assay, and those with a negative DNA methylation test result are spared a repeat biopsy, thereby reducing the number of unnecessary procedures.

The cost analysis was based on total costs of 1 year in the course of prostate cancer screening and evaluation through prostate biopsy. Costs to the health plan are assumed to be equal to the Medicare FFS rates, which provide a conservative benchmark for reimbursement rates paid for by other health plans.

Methods

Study Design

The budget impact model is presented from the perspective of a hypothetical commercial health plan, and direct costs are calculated over a 1-year time horizon, using Medicare fee-for-service (FFS) rates. The membership of this health plan is based on an assumed size of 1 million members, half of whom are males. The membership is distributed among age categories according to US population data.

The cost analysis was based on total costs of 1 year in the course of prostate cancer screening and evaluation through prostate biopsy. Costs to the health plan are assumed to be equal to the Medicare FFS rates, which provide a conservative benchmark for reimbursement rates paid for by other health plans.
scenario, the health plan incurs the costs of 1 or several repeated biopsies and the associated iatrogenic costs. In the new scenario, the health plan incurs the cost of an additional diagnostic test performed on the residual prostate tissue from the original sample plus the cost for repeated biopsies and associated iatrogenic costs on patients who had positive test results.

The model assumes that the epigenetic assay would be used for all men meeting the assay’s eligibility requirements, including an abnormal DRE, an elevated PSA level, and a negative prostate biopsy. Both scenarios calculate a total cost and a plan budget impact, expressed on a per-member per-month (PMPM) basis, as well as the aggregate annual cost to the plan.

Data Sources
The PubMed database was searched for published clinical and pharmaco-economic studies to assign values to the clinical and cost parameters used in the model. Studies were identified in PubMed that reflect current practice patterns of 10 to 12 core prostate biopsies in contrast to older studies that were based on sextant biopsy practice.

For cost parameters, a combination of published literature cross-referenced to Medicare payment rates was used. Parameters pertaining to the accuracy and outcomes of the assay (including the assay’s sensitivity and specificity, and the positive and negative predictive values) were cited from the MATLOC clinical trial.8

Sample Selection
PSA screening rates in US males vary by age, ranging from approximately 8% to almost 50%.24 Of all men screened for PSA, 6.8% are assumed to undergo biopsy based on a PSA of at least 4 ng/mL.25 These PSA and biopsy rates were applied to the hypothetical commercial health plan. Of the hypothetical plan, patients aged 40 to 64 years were selected, using demographic data from the US Census Bureau 2010, for inclusion in the model. Applying the national PSA screening and biopsy rates to the hypothetical commercial population yields a total of 2801 men undergoing a prostate biopsy. A total of 2101 men were deemed at risk for repeated biopsy based on a prostate cancer detection rate of 25% (Figure).25

Total Men at Risk for a Repeated Biopsy
This budget impact analysis compares the proportion of the 2101 men in the cohort who are at risk for a repeated biopsy in 2 potential scenarios—the reference scenario and the new scenario, which is using the epigenetic assay.

In the reference scenario, many patients with rising or elevated PSA levels will be seen again for DRE or PSA testing, and be considered for a repeated biopsy. In this standard of care, 43% (903) of the patients with a histopathologically negative biopsy are referred for a repeated biopsy. In this scenario, 43% (903) of the patients with a histopathologically negative biopsy are referred for a repeated biopsy based on persistent clinical risk factors.13

In the new scenario, these same patients are triaged with the epigenetic assay. In the new scenario, 3% of patients would not be eligible for the epigenetic assay, because of atypical small acinar proliferation (ASAP) found in their previous biopsy tissue.12 The model assumes that patients with ASAP will receive a repeated biopsy, given the high risk of prostate cancer associated with this histopathologic finding. Approximately 99.9% of all cases
would have sufficient tissue for this epigenetic assay, leaving 875 evaluable cases (based on 2 quality-not-sufficient cases of 749 in laboratory experience, through November 30, 2012, when the data were collected).

The Epigenetic Assay

The multiplex DNA methylation epigenetic assay became available in the United States in May 2012 (through MDxHealth’s CLIA-accredited, CAP-certified laboratory in Irvine, CA). The performance characteristics of this assay (Table 1) were described in the MATLOC clinical trial, which investigated the clinical utility of this assay.8 These characteristics were used in our budget impact model to determine the anticipated number of patients who would be identified as positive or negative for methylation markers.

The test is designed such that its high (90%) negative predictive value accurately distinguishes the patients with negative prostate biopsies from patients who may have occult cancer.8 In the budget impact model, under the reference scenario, 43% (903) of men at risk for repeated biopsy were referred for repeated biopsy. In the new scenario, testing the high-risk patients with the epigenetic assay significantly reduced the number of repeated biopsies by confirming the histopathologically negative biopsy results for 510 men. The epigenetic assay identified 365 men with positive DNA methylation results who would be referred for repeated biopsy (Table 2).

Model Variables and Assumptions

Conservative assumptions were made as to the number of repeated biopsies based on reported rates. In the standard of care, 43% of men are referred to have 1 repeated biopsy, 44% of which have a second biopsy, and 43% of these have a third biopsy. The 100% of men tested with the epigenetic assay who are methylation-positive were assumed to receive a repeated biopsy. A total of 1472 repeated biopsies are expected to be performed in the reference scenario compared with only 365 repeated biopsies with the epigenetic assay in the new scenario.

The average cost of a prostate biopsy procedure is $1946, which is a conservative estimate, based on decreased interim 2013 Medicare Physician Fee Schedule rates; this does not take into account prophylactic antibiotic, pain, or other concomitant medication costs.25

The total expected complication costs per patient for an initial or repeated biopsy were calculated using Surveillance, Table 2

<table>
<thead>
<tr>
<th>Variable</th>
<th>Reference scenario (standard of care)</th>
<th>New scenario (epigenetic assay)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Men at risk of repeated biopsy, N</td>
<td>2101</td>
<td>2101</td>
</tr>
<tr>
<td>Men referred for repeated biopsy or methylation test, %</td>
<td>43</td>
<td>43</td>
</tr>
<tr>
<td>Number of men referred for repeat biopsy or methylation test, N</td>
<td>903</td>
<td>903</td>
</tr>
<tr>
<td>Number of men with ASAP, %</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Cases with sufficient tissue for methylation test, %</td>
<td>N/A</td>
<td>99.9%</td>
</tr>
<tr>
<td>Risk stratification based on methylation markers</td>
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<td></td>
</tr>
<tr>
<td>Evaluable cases, N</td>
<td>N/A</td>
<td>875</td>
</tr>
<tr>
<td>Cases with positive methylation markers, N</td>
<td>N/A</td>
<td>365</td>
</tr>
<tr>
<td>Cases with negative methylation markers, N</td>
<td>N/A</td>
<td>510</td>
</tr>
<tr>
<td>Total men referred for repeated biopsy, N</td>
<td>903</td>
<td>365</td>
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</table>

ASAP indicates atypical small acinar proliferation; N/A, not applicable.
Epidemiology and End Results (SEER)-Medicare cancer registries’ reported incidence of infectious and noninfectious complications and the associated mean payment for the Medicare Severity-Diagnosis Related Groups (MS-DRGs) from the 2012 Centers for Medicare & Medicaid Services MS-DRG payment schedule.22

Table 3 shows the calculations for the average cost ($392) of complications per patient undergoing repeated prostate cancer biopsy. The total weighted cost of a fully burdened biopsy is $1946—the sum of the procedural cost and the cost of complications weighted by incidence. The retail price for the epigenetic assay is $206 per individual core, or $2061 for a 10-core biopsy (pricing is based on the cost of the ConfirmMDx for Prostate Cancer Test, as provided by MDxHealth, the manufacturer of this test). The model assesses the health plan’s costs compared with billed charges; therefore, the cost of this assay is discounted by 10%, to conservatively reflect payer costs, at $1855.02 per test.

Results

Costs

This budget impact analysis demonstrates that the net cost to a commercial plan is lower if patients undergoing prostate cancer biopsies are managed using the assay. Although this involves an additional cost for the acquisition of the assay, using the assay results in a reduction of 1106 unnecessary biopsies for a health plan with 1 million members (Table 4).

The total cost of repeated biopsies avoided is $2,152,276 (1106 biopsies avoided × $1946 per biopsy). The total cost to the health plan in 1 year was calculated to be $2,864,142 in the reference scenario versus $2,333,341 with the epigenetic assay in the new scenario. To calculate the total diagnostic cost per patient in the reference scenario, the cost of a prostate biopsy ($1946) was applied and weighted to a repeated biopsy distribution rate for the percentage of men who receive first, second, and third repeated biopsies of 43%, 44%, and 43%, respectively.13

For the new scenario, the total diagnostic cost per patient includes the cost of the assay plus the weighted biopsy cost, applied and weighted to 43% of men who have positive results based on the epigenetic assay (based on the sensitivity, specificity, and negative and positive predictive values).8 The total diagnostic cost per patient was $3172 in the reference scenario compared with $2584 in the new scenario, resulting in a savings of $588 per patient managed. This results in a total savings of $530,801 annually to the health plan, or –$0.0442 PMPM (Table 5).

Sensitivity Analysis

To test for uncertainty among the model parameters,
Epigenetic Assay: Screening for Prostate Cancer

Table 5  Total Annual Costs and Budget Impact: Reference Scenario versus New Scenario

<table>
<thead>
<tr>
<th>Cost</th>
<th>Reference scenario (standard of care)</th>
<th>New scenario (epigenetic assay)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Total annual cost, $</td>
<td>PMPM cost, $</td>
</tr>
<tr>
<td>Total cost of epigenetic assay</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total cost of repeated biopsies</td>
<td>2,864,142</td>
<td>0.33</td>
</tr>
<tr>
<td>Total cost of repeated biopsies avoided</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total diagnostic cost to health plan</td>
<td>2,864,142</td>
<td>0.24</td>
</tr>
<tr>
<td>Total diagnostic cost per patient</td>
<td>3172</td>
<td>0.24</td>
</tr>
<tr>
<td>Total budget impact to plan</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

PMPM indicates per member per month.

Table 6  Sensitivity Analysis: Total Costs and Budget Impact

<table>
<thead>
<tr>
<th>Cost</th>
<th>Reference scenario (standard of care)</th>
<th>New scenario (epigenetic assay)</th>
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<tbody>
<tr>
<td></td>
<td>Total annual cost, $</td>
<td>PMPM cost, $</td>
</tr>
<tr>
<td>Total cost of epigenetic assay</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total cost of repeated biopsies</td>
<td>9,429,097</td>
<td>1.09</td>
</tr>
<tr>
<td>Total cost of repeated biopsies avoided</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
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<tr>
<td>Total budget impact to plan</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

PMPM indicates per member per month.

all calculations were repeated for a patient population of men aged ≥65 years, representative of the Medicare population. Patients older than age 74 years were excluded from the model, resulting in an at-risk cohort of 6917 men. Testing these patients with the assay resulted in a reduction of 3642 unnecessary biopsies. The total cost of repeated biopsies avoided was $7,086,416. The total cost to the plan in 1 year was $9,429,097 in the reference scenario and $7,688,849 in the new scenario. This resulted in a total budget impact of −$1,740,248 to the plan, or −$0.1450 PMPM. The total diagnostic cost per patient was $3172 in the reference scenario compared with $2584 in the new scenario (Table 6), resulting in a savings of $588 per patient managed.

Discussion
The budget impact model was developed to evaluate the clinical and financial benefits of payer coverage for the epigenetic assay, also considering the clinical benefits, based on well-founded and conservative assumptions from existing evidence and current standards of care for patients considered at risk for prostate cancer.

The analysis demonstrates that a commercial health plan would realize cost-savings with the coverage of the epigenetic assay. The upfront cost of the epigenetic assay will be recovered based on the savings associated with avoided biopsy procedures and associated complications of biopsies. Given these assumptions and the costs associated with the current standard of care, the inclusion of
the epigenetic assay into the management of men who are screened for prostate cancer would result in a net cost-savings of ~$330,801 in the first year after the assay became available in the United States in a health plan with 1 million members.

Approximately $1.8 billion is spent annually on PSA testing alone, and more than $4 billion is spent on prostate cancer therapies, leading the AUA to call for new biomarkers to improve accurate diagnosis and reduce the cost burden.

The epigenetic assay provides clear and actionable results that aid the urologist in treatment decision-making, improving patient care, and yielding significant healthcare savings. A key assumption is that a health plan inclusion of the epigenetic assay in medical policy and coverage decisions will motivate a change in the behavior of urologists, resulting in a reduction of repeated biopsies. Policy tools that promote appropriate patient management according to evidence-based guidelines, such as value-based payment (eg, financial incentives for choosing evidence-based interventions) or coverage restrictions for repeated biopsies, may further enhance such outcomes.

Limitations

The results of this budget impact analysis are based on a hypothetical cohort modeled on the basis of values from the published literature. The use of national averages may not reflect the true variety in clinical practice. Initial prostate biopsy and repeated biopsy rates in specific geographic regions may be higher or lower than the reported national averages. Costs and resource utilization may also vary between practices and between geographic regions.

Another potential limitation to the application of this model is that the future rates of screening for prostate cancer may vary, given the recent recommendation of the USPSTF to stop routine PSA-based prostate cancer screening. The recommendation suggests that physicians discuss the benefit-risk ratio with their patients and decide if PSA testing is appropriate based on risk factors such as race or family history. How this will affect screening rates is not yet known and is not explored in this analysis.

This study is intended to address the financial impact of the epigenetic assay on the costs to commercial health plans of repeated biopsies. Because the cost impact is associated with a reduction in complications from the biopsy, the study provides some perspective on the impact of the assay on clinical outcomes. However, clinical outcomes were not evaluated in the design of the present study. A cost-effectiveness analysis would be the suitable approach to investigate the cost and the clinical outcomes associated with the use of the assay.

In addition, the assay’s impact on the rates of prostate cancer diagnosis and earlier case identification were outside the scope of the study and were not methodically investigated.

Subgroup analyses were not performed for modestly elevated PSA patients versus those with markedly elevated PSA, because these patients are not managed differently in clinical practice.

Because of the negative predictive value of 90% of the epigenetic assay, 10% of patients testing negative with the assay could have a diagnosis of prostate cancer identified through recommended return screening. The clinical impact of this is not evaluated by the design of this model. The test has only been commercially available in the United States since May 2012.

Conclusion

Approximately $1.8 billion is spent annually on PSA testing alone, and more than $4 billion is spent on prostate cancer therapies, leading the AUA to call for new biomarkers to improve accurate diagnosis and reduce the cost burden. Epigenetic assays have been reported to improve the accuracy of prostate biopsies and help to prevent repeated biopsies, the majority of which show negative results. Based on a recent analysis, the results of a new epigenetic assay can guide urologists in decisions regarding the need to repeat a biopsy in patients with a previously negative biopsy who are still considered at risk for prostate cancer.

A budget impact analysis calculated whether this assay can also produce financial benefits, beyond the previously reported clinical benefits. Using a hypothetical health plan with 1 million members, this analysis shows that the total annual cost to the health plan would be reduced if patients with histopathologically negative biopsies would be managed with the epigenetic assay.

Specifically, the use of the assay would reduce the number of repeated biopsies from 1472 to 365, thereby preventing 1106 unnecessary biopsies and reducing the annual costs by approximately $500,000 to the health plan, based on the current standard of care. This test has only been available for a short time. Research to investigate the clinical impact of this essay based on real-world data will be appropriate.

Disclaimer

At the time this study was submitted for publication,
prostate biopsy costs incorporated maximum payment reductions; conversion factor $25,000.8 per the 2013 Medicare Physician Fee Schedule published December 5, 2012. If changes to the sustainable growth rate patch and sequestration cuts occur, savings for payers reimbursing the epigenetic assay may be greater than reported.

Author Disclosure Statement

Dr Aubry is a Consultant to MDxHealth. Dr Lieberthal receives research/grant support from Abbott Molecular, Genomic Health, and MDxHealth. Dr A. Willis is a Consultant to ConfirmMDx and 21st Century Oncology, and is on the Speaker’s Bureau for Astellas, Clinlogix, GlaxoSmithKline, Eli Lilly, and sanofi aventis. Dr Bagley is a Consultant/Advisor to HillCo HEALTH. Mr Layton is a Consultant to ConfirmMDx. Mr S.M. Willis has nothing to disclose.

References


STAKEHOLDER PERSPECTIVE

Molecular Epigenetic Tests Can Improve Clinical Outcomes While Reducing Healthcare Costs

By Kelly Huang, PhD
President, HealthTronics, Inc, Austin, TX

HEALTH PLANS: Health insurance plans and other payers recognize the potential for molecular diagnostics to facilitate the approach known as personalized medicine, which utilizes molecular testing to identify patients who will benefit from a specific approach to management or a specific targeted therapy. Personalized medicine can lead to reduced healthcare costs over the life of the patient. However, payers are struggling to keep up with the rapidly expanding range of molecular tests.

Because of the ambiguous nature of “laboratory-developed tests,” many health insurers are deferring many new tests as experimental or investigational and are therefore refusing to cover them. When determining coverage of molecular diagnostic tests, payers expect evidence not only for clinical utility but also for cost-effectiveness. It follows that cost-effectiveness should also lead to a reimbursement structure that is based on the value of that test or service rather than merely on stacking codes of the methodology steps.

In this current article by Aubry and colleagues, the
authors propose a well-articulated hypothetical model to assess the value of an epigenetic test’s (ConfirmMDx) ability to confirm the negative results of a prostate biopsy. According to this model, using this test leads to a meaningful $588 savings per patient managed, by avoiding unnecessary repeated prostate biopsies.

As with any hypothetical model, there are limitations to the analysis with regard to real-world facts. Nevertheless, the authors clearly outline their assumptions and the rationale behind their calculations, so that readers can consider the conclusions, assess the sensitivity of the assumptions, and arrive at their own perspective.

Beyond the economic benefits of this test, payers should also consider the improvement in quality of life for patients who avoid the need for repeated biopsies.

**Patients/Providers:** Patients and providers will experience the benefits of many new and soon-to-become-available advances in the detection and treatment of prostate cancer. Along with new drug therapies and surgical advances, such as minimally invasive ablative procedures, molecular epigenetic tests show promise in determining the aggressive nature of a tumor, or in the case of ConfirmMDx, provide true confirmation of negative biopsy results.

Although these epigenetic tests are relatively new, they already provide further input, along with the details of the biopsy results, prostate-specific antigen history, family history, digital rectal examination, age, and so on, to help the physician’s and the patient’s determination of a specific treatment regimen. Over time, as more experience is gained and the tests are improved for specificity and sensitivity, molecular epigenetic tests can be expected to provide significant improvements in extended survival and enhanced quality of life for patients.


Disclosure: HealthTronics offers Laboratory Solutions, including ConfirmMDx and other epigenetic testing for prostate cancer.