



Px SCORE[®]
25
LOW RISK

POST-OP[®] Px
THE POWER TO PREDICT

AFTER PROSTATE
CANCER SURGERY...

Find out your
Px SCORE[®].

AUREON[®]
BIOSCIENCES



Introducing Post-Op Px[®]

Now there's a personalized, objective, and highly accurate test for patients post-surgery.

Q. *Why should I consider Post-Op Px[®] after I've had my prostate removed?*

A. Many physicians consider surgery the gold standard and it is an effective treatment. However, post-surgery, a number of men have adverse, higher risk features such as positive surgical margins, extracapsular extension, seminal vesicle invasion or even higher than expected Gleason scores, which can be a source of anxiety for men and their loved ones.

Current assessment of these high-risk features (e.g., positive surgical margins) is challenging and subjective.

If you have concerns after surgery, Post-Op Px provides an objective and more accurate perspective about your likelihood of cancer recurrence.



Q: *How Accurate is Post-Op Px®?*

A: If Post-Op Px identifies you as low risk for clinically significant disease, then there is a 99% likelihood that you will not fail salvage therapy or experience metastasis within five (5) years after surgery.*

Q: *Is Post-Op Px right for you?*

A: If you have had surgery to remove your prostate within the last five (5) years, please answer the following questions with your doctor to see if you are a good candidate for Post-Op Px:

1. *Did your prostatectomy tissue have positive surgical margins (SM)?* Yes No
2. *Did your prostatectomy tissue have extracapsular extension (ECE) or seminal vesicle invasion (SVI)?* Yes No
3. *Did your prostatectomy tissue have a higher Gleason result?* Yes No
4. *Has your PSA come back (after surgery)?* Yes No
5. *Do you have any anxiety about the cancer recurring?* Yes No

If you answered “Yes” to any of the above questions, please ask your doctor about Post-Op Px or call us for more information that you can share with your physician.

* As assessed in the validation cohort of the Post-Op Px developmental study.

Q: *What is Post-Op Px[®] and how does it work?*

A: Post-Op Px is a test that is ordered by your doctor. No additional laboratory visits or needle sticks are required.

When Post-Op Px is ordered, a piece of your prostate tissue (from your surgery) is collected from the pathology department of the hospital where your surgery was performed and sent to our specialized laboratory for further analysis.

Your samples will be put through a rigorous process using state-of-the-art technology to scan your tumor for specific cellular and molecular patterns that help predict cancer aggressiveness. This new analysis will help you and your physician to better understand your risk for having your disease return as well as your risk for clinically significant disease.



Q: *What makes Post-Op Px® so different?*

A: In a word... technology.

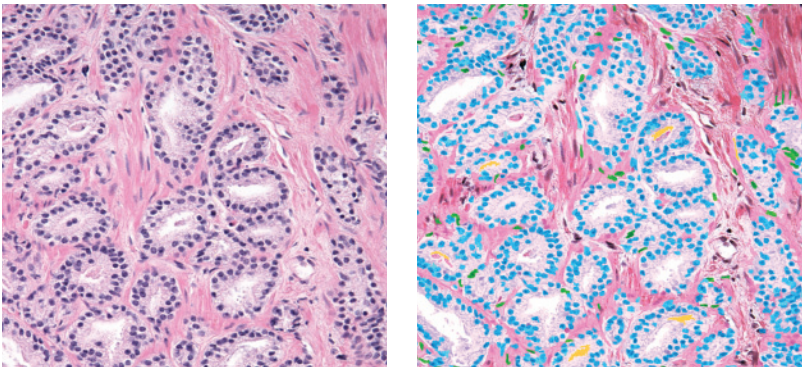
Thanks to its breakthrough, patented technology, Post-Op Px is the first and only test that combines information from multiple domains to predict the return of clinically significant cancer.

Cellular: Post-Op Px takes a detailed look at the cells that make up the tumor.

Molecular: Post-Op Px is able to evaluate the proteins in your prostate tissue to get a unique biologic picture of what's happening.

Clinical: Post-Op Px takes into account important clinical information such as Gleason score, PSA levels, and your surgery findings.

By combining this information and using advanced, one-of-a-kind computer technology, Post-Op Px is able to deliver a thorough and accurate picture of your individual risk.



The above images illustrate just one of many steps that tissue must undergo for Post-Op Px. The image at the left was taken before Post-Op Px analysis. The image at the right shows the same tissue after analysis.

Q: What will my doctor and I receive once the test is done?

A: Your doctor will receive a report with your individualized test results and will share the findings with you.

The report includes your Px SCORE® which tells you the likelihood of your prostate disease returning and if it will be clinically significant. Your Px SCORE can range from a low of 0 to a high of 100. The lower the score, the lower the likelihood that your disease will return.



Q: What happens after my doctor receives the Post-Op Px® report along with my Px SCORE®?

A: Your doctor will use your Px SCORE, in combination with additional information such as Gleason score and PSA level, to draw a more complete and objective picture of your future health and provide more-informed counseling. Getting an accurate prediction of the future can help relieve the stress and anxiety surrounding the unknown and allow you, your family, and your doctor to decide the best path forward.



Q: *What about Reimbursement?*

A: Aureon currently bills all commercial, private, third party carriers, and Medicare. Our Aureon Patient Care Program – APCP can be helpful to patients in understanding the reimbursement process. Aureon feels very strongly that no patient should ever be penalized financially while working through their battle with Prostate Cancer.

Q: *How do I learn more?*

A: To find out more and to determine if the test is right for you, please speak with your doctor.



POST-OP[®] Rx
THE POWER TO PREDICT

For more information, visit www.aureon.com
or call **1-888-SYS-PATH** (797-7284).

Find out what your future holds – with Post-Op Px®

Post-Op Px:

- Can help reduce the anxiety associated with high-risk features (e.g., positive surgical margins) after surgery.
- Can tell you the likelihood of your prostate disease returning following prostate surgery.
- Will help you and your doctor to make future treatment decisions.
- Is easy to order through your doctor and requires no additional patient procedures.

For more information, visit www.aureon.com
or call **1-888-SYS-PATH (797-7284)**.



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The test was developed and its performance characteristics determined by Aureon Biosciences, Inc. (Aureon). Clearance or approval of the U.S. Food and Drug Administration was not required for this test as of the date of this publication. Aureon is regulated under the Clinical Laboratory Improvement Amendments of 1988 (CLIA) and is qualified to perform high-complexity clinical testing. This test can be used for clinical purposes. These results are adjunctive to the ordering physician's workup.

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