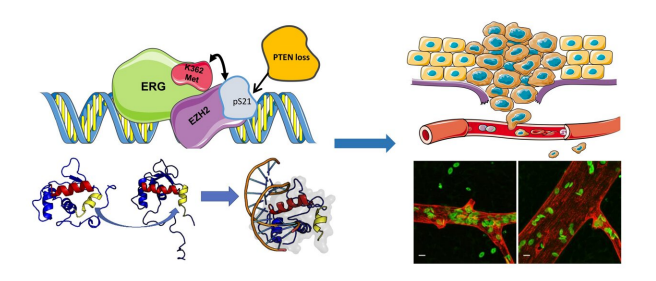


## A molecular switch turns on the unfavorable evolution of prostate tumors

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EZH2 induces ERG methylation at lysine 362 (K362), leading to a conformational change that enhances the transcription factor ERG's DNA binding and trans-activating capability. Phosphorylation of EZH2 at serine 21 (pS21) promoted by loss of PTEN increases the ability of EZH2 to methylate ERG. K362 methylated ERG causes a broad transcriptional reprogramming with activation of genes enhancing intravascular dissemination and metastatic capability promoting prostate cancer progression. Credit: Università della Svizzera italiana

Researchers from the Prostate Cancer Biology laboratory, directed by Giuseppina Carbone, M.D., at the Institute of Oncology Research (IOR, affiliated to USI) in Bellinzona, have discovered an unexpected



mechanism that drives the largest group of prostate tumors' evolution, the ERG fusion-positive prostate cancers. The study is published in *Nature Communications*.

Prostate <u>cancer</u> is one of the most common malignancies and causes of mortality in men worldwide. About 50% of <u>prostate</u> tumors (ERG fusionpositive prostate cancers) harbor a fusion between the ERG gene and the promoter region of TMPRSS2 gene. This chromosomal rearrangement leads to abnormal production of ERG and promotes tumor progression. However, the mechanism by which ERG contributes to the tumorigenesis and drives the progression from primary to metastatic and hormone-refractory prostate cancers is still unclear.

Looking for proteins that potentially interact and cooperate with ERG, researchers identified EZH2. Dr. Carbone explains: "We discovered that EZH2 binds and adds a <u>methyl group</u> to ERG, acting as a co-activator. EZH2 is a partner-in-crime of ERG in prostate cancer progression."

The researchers observed that the addition of a methyl group on a specific site of ERG was enhancing its tumorigenic activity. "The methylation of lysine 362 is like a switch turning ERG from a quiescent to a very active state. This modification occurs in a critical position at the boundary of an internal auto-inhibitory domain in the ERG protein," says Dr. Cavalli.

With the methylation on lysine 362, ERG changes its conformation and increases its activity, promotig the expression of multiple genes involved in tumor progression and metastasis. Using preclinical models of ERG fusion-positive prostate cancer, the researchers unraveled the role of another important player in EZH2-ERG activation: the gene PTEN. Deletion of PTEN are widespread in prostate cancers and often associated with ERG fusion-positive tumors. Dr. Carbone's group observed that the loss of PTEN increased EZH2 activity and, therefore



ERG methylation, thus explaining the combined effects of these events in patients.

"Our findings define a new therapeutically actionable pathway for treating patients with ERG fusion-positive prostate cancer and preventing further progression of the disease," says Prof. Catapano. "In various experimental models, drugs that inhibit EZH2 and prevent ERG methylation reverse the molecular and phenotypic alterations induced by ERG and block <u>tumor</u> growth. This strategy could turn to be very effective in the more aggressive ERG fusion-positive prostate tumors. Interestingly, other tumors expressing ERG might exploit similar mechanisms of EZH2-mediated activation. These aspects are currently under investigation."

**More information:** Marita Zoma et al, EZH2-induced lysine K362 methylation enhances TMPRSS2-ERG oncogenic activity in prostate cancer, *Nature Communications* (2021). DOI: 10.1038/s41467-021-24380-6

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