PSP94, an upstream signaling mediator of prostasin found highly elevated in ovarian cancer

J-x Ma1,2,6, B-x Yan1,3,6, J Zhang3,4, B-H Jiang5, Y Guo1, H Riedel1, MD Mueller1, SC Remick1 and JJ Yu1

Ovarian cancer is a leading cause of cancer death as diagnosis is frequently delayed to an advanced stage. Effective biomarkers and screening strategies for early detection are urgently needed. In the current study, we identify PSP94 as a key upstream factor in mediating prostasin (a protein previously reported to be overexpressed in ovarian cancer) signaling that regulates prostasin expression and action in ovarian cancer cells. PSP94 is overexpressed in ovarian cancer cell lines and patients, and is significantly correlated with prostasin levels. Signaling pathway analysis demonstrated that both PSP94 and prostasin, as potential upstream regulators of the Lin28b/Let-7 pathway, regulate Lin28b and its downstream partner Let-7 in ovarian cancer cells. Expression of PSP94 and prostasin show a strong correlation with the expression levels of Lin28b/Let-7 in ovarian cancer patients. Thus, PSP94/prostasin axis appears to be linked to the Lin28b/Let-7 loop, a well-known signaling mechanism in oncogenesis in general that is also altered in ovarian cancer. The findings suggest that PSP94 and PSP94/prostasin axis are key factors and potential therapeutic targets or early biomarkers for ovarian cancer.

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Epithelial ovarian cancer is the most lethal gynecologic cancer and the fourth leading cause of death among women in developed countries. If diagnosed at stage I with the malignancy confined to the ovary, the 5-year survival rates can reach up to 94%. However, survival rates sharply decline to 28% for advanced stages at presentation.1 Insufficient biomarker(s) and the lack of an effective screening strategy for early detection result in >80% of patients presenting with advanced disease at the time of diagnosis. Serum marker CA125 (cancer antigen 125) is clinically used in ovarian cancer screening, but it lacks the sensitivity and specificity to function independently in a test.2 A sensitive biomarker for accurate early detection of ovarian cancer is urgently needed.

Prostasin (protease serine 8 or PRSS8) is a trypsin-like serine peptidase expressed in epithelial cells and is found as a cell surface bound or secreted protein. Prostasin is altered in ovarian, prostate, breast and gastric cancers3–6 and has been proposed to inhibit cancer cell proliferation and invasion upon activation.7 In particular, prostasin is overexpressed in ovarian cancer and may serve as a potential biomarker for early detection of ovarian cancer independently or in combination with CA125.4,8,9

Prostatic secretory protein 94 (PSP94) also termed microsemiprotein beta (MSMB) is the second most abundant protein in the semen of healthy men and is found in a variety of human tissues.10–12 The cellular levels of PSP94 gradually decreases with the progression of prostate cancer, suggesting that PSP94 may represent a promising target for cancer treatment and a potential biomarker for early detection of prostate cancer.13 In addition, genome-wide association studies and functional analysis of underlying MSMB gene correlate with polymorphism of the gene promoter region, resulting in altered MSMB expression with prostate cancer risk.14–17

MicroRNAs (miRNAs) regulate gene expression at post-transcriptional level in diversely biological functions such as cell proliferation, differentiation and apoptosis.18,19 The Let-7 family of miRNAs consists of more than 10 sequence-conserved members with similar functions across diverse species from worms to humans.20 Deregulation of Let-7 has been linked to many types of cancer and other diseases.20,21 An RNA-binding protein termed Lin28b was recently identified as a direct upstream inhibitor of Let-7 family signaling22,23 with important functions in embryonic stem cells and embryonal carcinoma cells.24–27 Balanced signaling of Lin28b to Let-7 is critical; imbalance is linked to various diseases.28

Our recent studies demonstrated important roles and functional similarity of prostasin and PSP94 in ovarian cancer chemoresistance,29,30 In the current study, we report that prostasin expression is regulated by PSP94 and shares all examined downstream targets with PSP94 in ovarian cancer cells. PSP94 and prostasin are both overexpressed in strong correlation with Lin28b/Let-7 expression levels in tissues of ovarian cancer patients. As a result, PSP94 and the
PSP94/prostasin axis may represent promising markers for early detection of ovarian cancer as well as potential therapeutic targets. This role may be extended to other types of cancers where either one or both components are altered.

Results

**PSP94 regulates prostasin expression in ovarian cancer cells.** PSP94 and prostasin both play important roles in chemoresistance and share similarities in their actions.\(^{29,30}\) To investigate their possible relationship, we examined PSP94 expression in prostasin overexpressing cells (O432-RP-pro-O) and prostasin level in PSP94 overexpressing cells (O432-RP-PSP-O). PSP94 expression was found unchanged in O432-RP-pro-O when compared to control cells (Figure 1, lane 3, top band), whereas prostasin level was found increased in O432-RP-PSP-O cells when compared to control cells O432-RP-C (Figure 1, lane 2, top band). These findings are consistent with a role of PSP94 as an upstream regulator for inducing prostasin expression, but not vice versa. We extended this observation to a different ovarian cancer cell line where, similarly, prostasin expression increased upon PSP94 overexpression (Supplementary Figure 1). We did not examine PSP94 expression since we were unable to produce prostasin overexpression, possibly due to the strong repression function of prostasin in ovarian cancer cells.\(^{30}\)

**Prostasin is a downstream mediator of PSP94 in a shared signaling pathway.** We previously showed that prostasin regulates downstream signaling of the CASP/PAK2-p34 and MLCK/actin, JNK/c-Jun pathways.\(^{30}\) PSP94 may share these pathways with prostasin as an upstream mediator. We examined expression levels of various genes in CASP/PAK2-p34 and MLCK/actin, JNK/c-Jun pathways in PSP94 overexpressed cells O432-RP-PSP-O. All respective genes were found to be altered in O432-RP-PSP-O cells when compared to the control O432-RP-C (Figure 2a), and showed the same pattern as in prostasin overexpressed cells (O432-RP-pro-O when compared to the control).\(^{30}\) Our findings suggest shared signaling mechanisms between PSP94 and prostasin in CASP/PAK2-p34 and MLCK/actin, JNK/c-Jun pathways. PSP94 has been shown to regulate the Lin28/Let-7 loop in ovarian cancer cells.\(^{29}\) To investigate whether prostasin participates in this mechanism together with PSP94, we examined Lin28b and Let-7 expression levels in prostasin overexpressed cells O432-RP-pro-O. Lin28b level was increased, whereas Let-7 miRNA level decreased in O432-RP-pro-O cells compared to the control (Figures 2b and c). Based on our observations, PSP94 appears to function as an upstream mediator of prostasin and both proteins share all tested downstream targets. PSP94 regulates prostasin expression, but prostasin does not appear to affect PSP94 level.

PSP94 and prostasin have similar roles in chemoresistance in ovarian cancer\(^{29,30}\) and share downstream targets. Our data suggest a role for prostasin as a key downstream target of PSP94 in the development of chemoresistance. To investigate this hypothesis, we studied O432-RP-PSP-O-pro-D cells obtained by prostasin-siRNA transfection of O432-RP-PSP-O cells (Figure 2d). Both cell lines were treated with paclitaxel and cell survival was evaluated. As shown in Figure 2e, the survival of O432-RP-PSP-O-pro-D cells increased in a dose-dependent manner that indicates greater resistance to paclitaxel treatment when compared to O432-RP-PSP-O cells. Knockdown prostasin in O432-RP-PSP-O cells appeared to interfere with PSP94 action in chemoresistance by disconnecting PSP94 from its downstream targets. These data demonstrate that PSP94 is dependent on prostasin function in the development of chemoresistance in our experimental model.

**PSP94 or the PSP94/prostasin axis is overexpressed in ovarian cancer cell lines and ovarian cancer patients.** Prostasin is overexpressed in ovarian cancer patients and several ovarian cancer cell lines\(^4\) and PSP94 show a comparable expression pattern. When we compared PSP94 protein levels in normal and several ovarian cancer cell lines, we found high or modest levels of PSP94 expression in all cancer cell lines but not detectable in normal ovarian cells (Figure 3a). We also quantified PSP94 and prostasin mRNA expression in ovarian cancer tissues. As shown in Figure 3b, PSP94 mRNA levels are significantly higher (about four-fold, on average) in tumor tissues of ovarian cancer patients (at all stages) than in normal specimens. We also compared mRNA levels in early and late stages of the disease (stages 1 and 2 were defined as early stage, versus stages 3, 4 and 5 as late stage). We observed that PSP94 levels are four to five-fold higher in early stage samples and slightly elevated above control level in the late stages. A similar pattern of prostasin expression was also observed in these samples (Supplementary Figure 2).

Correlation analysis of PSP94 and prostasin expression showed significant overlap in ovarian cancer tissues (Figure 3c). Among 13 matched samples from ovarian cancer...
patients (tumor tissue and normal ovary tissue from the same patient) 11 were seen PSP94 expression increased in tumors compared to normal tissue (\( B^85\%\); \( \Delta^c^{43.6} \) value considered as positive expression and vice versa; Figure 3d).

PSP94 protein is secreted in various human tissues including breast and ovaries. 12 To evaluate whether PSP94 could serve as a serum biomarker for early detection of ovarian cancer, we examined PSP94 protein level in the blood of normal and ovarian cancer patients. In ovarian cancer patients (mixed stages), PSP94 levels were \( 35.7 \pm 21.7 \) ng/ml, which is significantly higher compared to normal controls \( (14.6 \pm 12.7 \) ng/ml; Table 1; \( P<0.001 \)), confirming that PSP94 is overexpressed in ovarian cancer patients.

PSP94 and prostasin regulate the Lin28/Let-7 loop in ovarian cancer cells and show correlation with Lin28/Let-7 expression in cancer patients. Expression of Lin28b and Let-7 is altered in ovarian cancer. 31,32 Our previous

Figure 2 Prostasin is a downstream mediator of PSP94 in a shared signaling pathway. (a) CASP2-p34 signaling and further downstream JNK/c-Jun and MLCK/actin signaling are regulated by PSP94. Expression levels of CASPs, Pak2-p34, JNK, c-Jun, MLCK and actin were examined in PSP94 overexpressed O432-RP-PSP-O cells, which show the same pattern as prostasin overexpressed cells O432-RP-pro-O. Overexpression of prostasin in O432-RP-pro-O cells upregulates CASP/Pak2-p34 and further downstream JNK/c-Jun and MLCK/actin signaling molecules (lanes 1, 2 and 3) as previously reported, 30 indicating that PSP94 and prostasin share the same downstream targets. (b and c) Prostasin regulates Lin28b/Let-7 loop in ovarian cancer cells. Expression levels of Lin28b and Let-7 microRNA were examined in prostasin overexpressed cells (O432-RP-pro-O) and the same expression pattern is seen as in PSP94 overexpressed O432-RP-PSP-O cells. Upregulation of Lin28b by overexpression of PSP94 (O432-RP-PSP-O; lanes 1, 2 and 4) is consistent with our previous report, 29 indicating that PSP94 and prostasin share the same downstream targets. (d) Prostasin-siRNA knockdown. Prostasin protein levels are decreased in O432-RP-PSP-O-pro-D knockdown cells compared to O432-RP-PSP-O (mock transfection control) or O432-RP-PSP-O-Cs (siRNA transfection control). (e) Prostasin-siRNA knockdown in O432-RP-PSP-O cells appeared to interfere with PSP94 action in chemoresistance. Cells were plated at 10–20% confluence, treated with paclitaxel at different concentrations for 24 h, and then cultured in normal medium for 7–10 days. Cell survival was evaluated. Relative survival rates of each cell line are shown.
Cell Death and Disease

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is a possible biomarker for early detection of ovarian cancer. 

Other studies report that prostasin is overexpressed in 

ovarian cancer and in combination with CA125 provides high 
sensitivity (92%) and specificity (94%) to identify cancer 

patients, but low sensitivity when used as an individual 

marker. We propose PSP94 as a more advantageous 

biomarker for earlier detection of ovarian cancer, compared 
to prostasin. PSP94 is a secreted protein, whereas 

prostasin is a secreted and membrane protein (glycosylpho-

phatidylinositol-anchored), with functional differences 
between the two proteins. 10,30 Prostasin appears to be 

implicated in a much wider spectrum of physiological and 

pathophysiological conditions than PSP94 and has important 
roles in epidermal barrier function, skin phenotype, embryonic 
viability and blood pressure. 34–38 In contrast, known functions 
of PSP94 are relatively narrow in prostate cancer develop-
ment and ovarian cancer chemo resistance. Although PSP94 
is a promising biomarker for ovarian cancer detection and 
treatment, further studies are needed. Large patient samples 
are required to determine sensitivity and specificity and 
functional studies are necessary to investigate PSP94 (and 
its peptide derivative PCK3145) as potential therapeutic 
targets. 

Our findings show that PSP94 is an upstream mediator of 
prostasin in ovarian cancer. PSP94 and prostasin are over-
expressed in ovarian cancer patients, and PSP94/prostasin 
signaling may play an essential role in ovarian cancer 

Discussion

Despite dramatic advances in recent decades in understand-
ing basic cancer mechanisms, ovarian cancer is still a 
deathy disease with a five-year survival rate of <50%. In 
particular, the lack of effective biomarkers for early screening 
results in >80% of patients presenting with advanced disease 
at the time of diagnosis. In this study, we show that PSP94 is 
overexpressed in ovarian cancer patients and cell lines and 
functions as an upstream mediator of prostasin. Since PSP94 
expression is particularly elevated in early-stage patients 
(stages1 and 2), and only somewhat in late-stage patients 

Figure 3  PSP94 expression levels are elevated in ovarian cancer cells and tumor tissues. (a) PSP94 expression levels by immunoblotting with specific antibodies in human ovarian normal and cancer cell lines. PSP94 protein levels are shown in all ovarian cancer cell lines Ovca432, Ovca433, Ovcar3, Caov3 and 2008, but no PSP94 expression is seen in normal ovarian surface epithelial (HIO-80) cells. (b) PSP94 was overexpressed in ovarian cancer patients. PSP94 mRNA levels were measured in tumor tissues of ovarian cancer patients and compared among different stages by real-time qPCR analysis. Relative levels of PSP94 in patients (early, late, unknown and all stages) were compared to normal controls. In this study, early stage n = 43, late stage n = 14, unknown stage n = 19 and normal control, n = 35. (Stages 1 and 2 were defined as early stage and stages 3, 4 and 5 as late stage.) P < 0.05. Mean ± S.D. are given, and the P-values were calculated using the two-sided Student’s t-test. (c) Correlation between PSP94 and prostasin in ovarian cancer patients. (d) Comparison of PSP94 levels in 13 patients with matched ovarian tumor and normal tissues. PSP94 mRNA levels were examined by real-time qPCR analysis. The Ct value of each matched samples (normal and tumor tissues of same patient) was compared. Ct = 36 was set as non-detectable level
development. PSP94 and prostasin levels are both decreased in prostate cancer; this is consistent with our preliminary findings that PSP94 regulates prostasin expression in prostate cancer cells and likely serves as an upstream signaling mediator of prostasin in prostate cancer development. We expect that PSP94/prostasin signaling may be important in oncogenesis in general, not just limited to ovarian cancer (Figure 5), and plays an important role in ovarian cancer chemoresistance. 29,30 Notably, PSP94/prostasin signaling appears to have an opposite role in ovarian cancer compared to prostate cancer, as PSP94 and prostasin are overexpressed in ovarian cancer,4 while their expression levels are decreased in prostate cancer.1,13 Prostasin expression is also altered in breast and gastric cancers5,6 and it may be meaningful to investigate the corresponding PSP94 expression in those cancers.

In our study, PSP94 is an upstream signaling mediator of prostasin in ovarian cancer cells, and both share all investigated downstream targets; however, prostasin may have other upstream regulators and downstream signals that are independent of PSP94 (Figure 5). This is suggested by some functional differences that we observed between PSP94 and prostasin in ovarian cancer cells. For example, recombinant PSP94 or PSP94-derived peptide PCK3145 represses some ovarian cancer cells, in contrast to recombinant prostasin. Overexpression of prostasin in ovarian cancer cells greatly induces cell death and represses cell survival, whereas overexpression of PSP94 in ovarian cancer cells does not significantly affect cell death.29,30 Thus, further studies on differences between these two genes in ovarian cancer are needed to comprehensively understand their role in oncogenesis.

Lin28b was recently identified as an RNA-binding protein and direct upstream mediator and inhibitor of Let-7 family.22,23 Alteration of Let-7 or Lin28b has been linked to many types of cancers and other diseases, and the Lin28/Let-7 loop is seen as a crucial signaling circuit in oncogenesis.20,21,32,39–41 Particularly, upregulation of Lin28b and altered Let-7 are associated with advanced disease and reduced patient survival in ovarian cancer25,31,43 and a Lin28b polymorphism has been linked to susceptibility to epithelial ovarian cancer.32 Our findings show that PSP94 and prostasin regulate Lin28b and Let-7 expression in ovarian cancer cells and their expression appear to be strongly correlated in ovarian cancer patients (Figure 4). Our data suggest that alteration of Lin28b and Let-7 expression in ovarian cancer may partially result from overexpression of PSP94 and prostasin. It is widely accepted that Lin28b overexpression contributes to tumorigenesis by repressing tumor suppressor Let-7 expression in ovarian and other cancers,20,21,32,39–41 which is consistent with our view that the PSP94/prostasin pathway may play a key role in the development of several cancer types (Figure 5). Our current findings suggest a potential link between Lin28b/Let-7 signaling and the PSP94/prostasin pathway in oncogenesis in general. The Lin28/Let-7 pathway is also believed to play a critical role in stem cell development.18,22,23 Thus, our new findings point to a
Cell survival assay. Cell survival was evaluated as previously described. 29 Cells were counted and plated into culture dishes at ~10–25% confluence on the day before treatment. Paclitaxel or PBS as control was added for 20 h and then removed. Cells were recovered and continuously propagated in normal medium for 10 days. To quantify final cell numbers, cells were stained with 0.25% crystal violet/20% ethanol and counted; or the proliferation rate was measured using a Cell Proliferation Assay kit (Promega, Madison, WI, USA) according to the manufacturer’s instruction. Briefly, MTS/PMS solution (at a final concentration of 333 μg/ml MTS and 25 μM PMS) was added to each well, and cells were incubated for 2–3 h at 37°C. The absorbance was determined at 490 nm using a 96-well plate ELISA reader. Culture medium was used as a background control. The experiments were repeated at least three times.

ELISA and PSP94 measurements. The plasma PSP94 concentration was determined using a PSP94 ELISA Kit (Sanbo Inc., Beijing, China); 100 μl of standard or plasma samples were loaded to 96-well plates and incubated for 1 h at 37°C with 50 μl of HRP conjugate. After rinsing, 50 μl of substrate A and 50 μl of substrate B were added to each well and incubated for 15 min at 37°C. 50 μl of Stop Solution (Sanbo Inc, Beijing, China) was added before the absorbance at 450 nm was determined with a 96-well plate ELISA reader.

Statistical analysis. The Student’s t-test and correlation analysis for Windows were used for statistical analysis with significance defined as P < 0.05. All statistical tests and corresponding P-values were two-sided.

Conflict of Interest
The authors declare no conflict of interest.
Supplementary Information accompanies this paper on Cell Death and Disease website (http://www.nature.com/cddis)