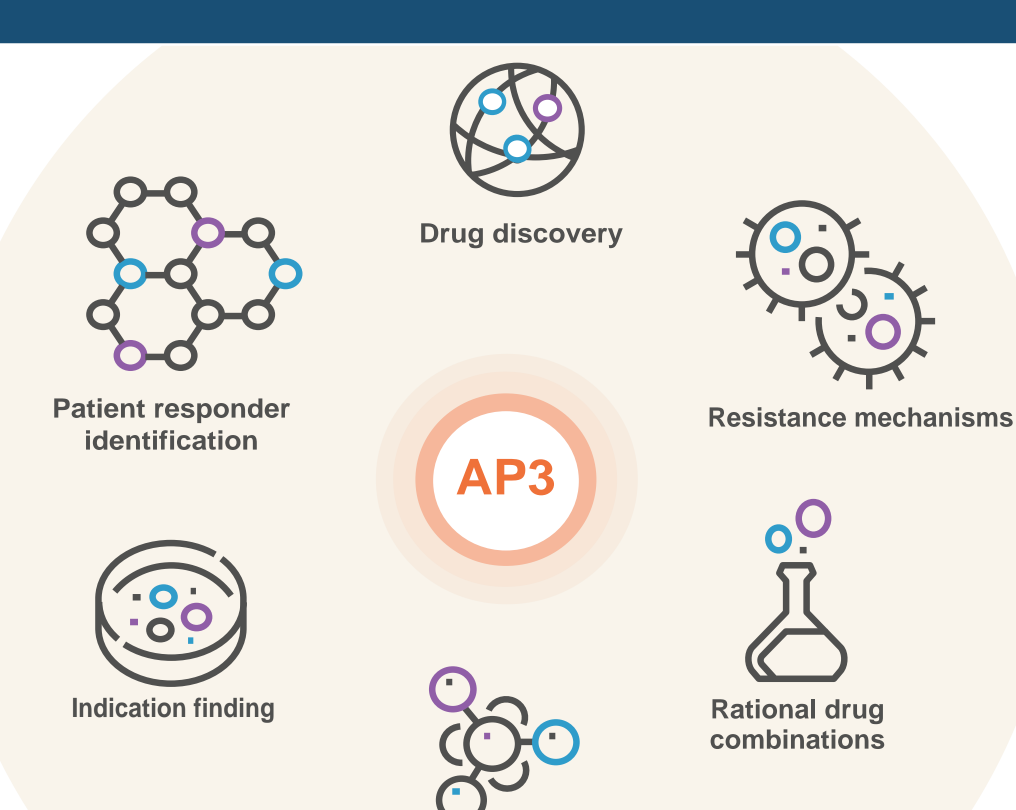
Acrivon Predictive Precision Proteomics (AP3) uncovers mechanism of resistance to ACR-368, a clinical-stage CHK1/2 inhibitor, and identifies rational combination treatment



Helén Nilsson¹, Lei Shi², Magnus E. Jakobsson¹, Joelle Baddour-Sousounis², Shahrzad Rafiei², Uthira Muralitharan¹, Zachary Best², Valentina Siino¹, Francisco J. Santana², Ignacio Arribas Diez¹, Kailash Singh², Ruban Cornelius¹, Nina Lipjankiç¹, Kate Rappard², Portia Lombardo², William Dahlberg², Subodh Kumar², Ahmed Youssef², Reina Improgo², Corey Xu², Joon Jung², Ung-Min Lee³, Ayesha Murshid², William Dahlberg², Subodh Kumar², Ahmed Youssef², Reina Improgo², Corey Xu², Joon Jung², Ung-Min Lee³, Ayesha Murshid², William Dahlberg², Subodh Kumar², Ahmed Youssef², Reina Improgo², Corey Xu², Joon Jung², Jung-Min Lee³, Ayesha Murshid², Michail Shipitsin², Jesper V. Olsen⁴, Kristina Masson¹, David A. Proia², Caroline Wigerup¹, Peter Blume-Jensen²

¹Acrivon AB, Medicon Village, Lund, Sweden, ²Acrivon Therapeutics, Watertown MA, USA, ⁴Center for Protein Research, University of Copenhagen, Copenhagen, Denmark

AP3 - Acrivon Predictive Precision Proteomics



Acrivon predictive precision proteomics (AP3) platform enables biological SAR based drug discovery and identification of drug-specific pharmacodynamic and response predictive protein biomarkers for drug indication screening and patient responder identification

In this study, the AP3 platform was used to uncover a key druggable resistance mechanism to ACR-368 and to identify a rational combination treatment to overcome resistance

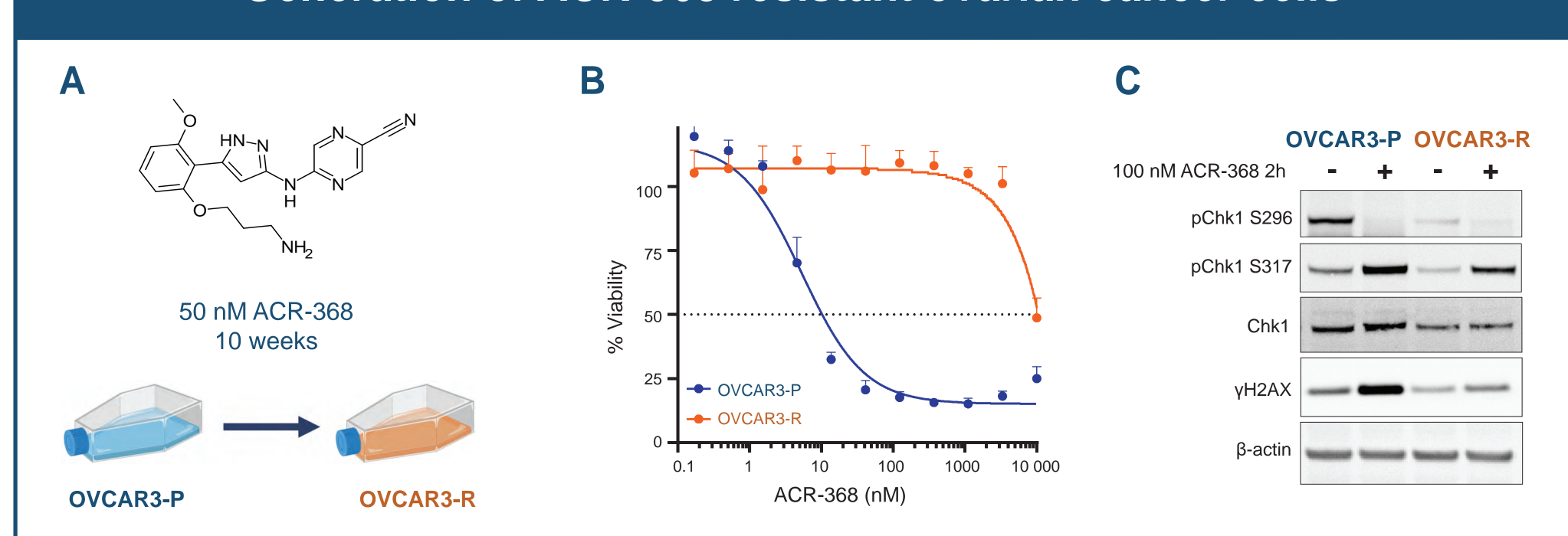
BACKGROUND

- ACR-368 is a potent and selective inhibitor of CHK1/2, key nodes in the DNA Damage Response (DDR) pathways, with demonstrated durable, single-agent activity in a subset of patients with advanced solid tumors
- As is the case for most DDR modulators, genomic biomarkers have also proven unsuccessful in predicting response to ACR-368, limiting the clinical success rates in patients
- Using the AP3 platform, we developed a drug tailored patient selection test ACR-368 Onco-Signature - for individualized patient drug response prediction
- A Phase 2 clinical trial is ongoing where patients are treated with ACR-368 monotherapy based on OncoSignature-predicted sensitivity (NCT05548296)

PRESENT STUDY

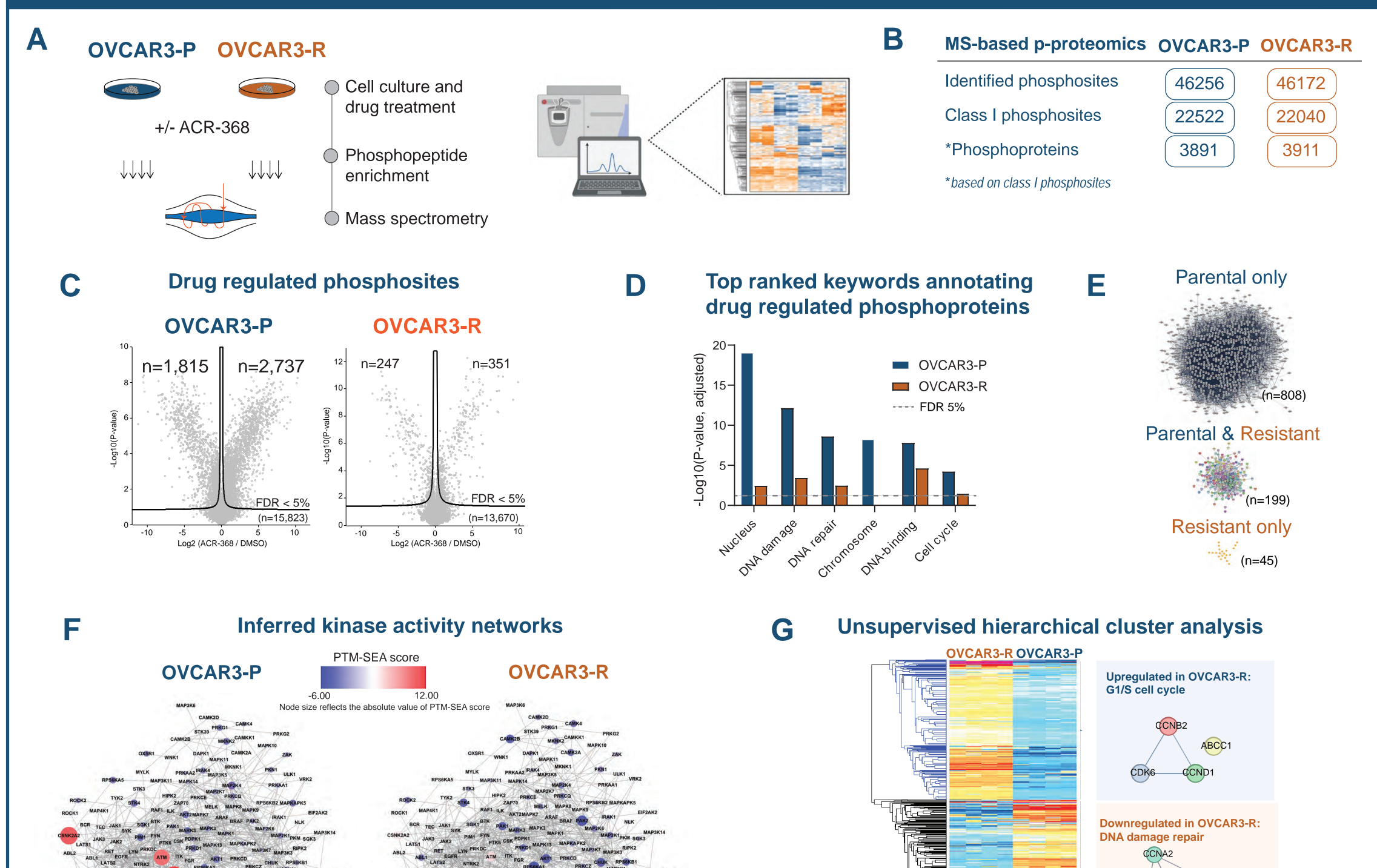
- > NIH-OVCAR3 (OVCAR3) ovarian cancer cells were generated durably resistant to ACR-368
- Differential response to ACR-368 treatment was profiled in OVCAR3 Parental (OVCAR3-P) and Resistant (OVCAR3-R) cells using AP3 mass spectrometry
- Comprehensive pathway reconstitution and kinase activity analysis was performed to identify drug resistance mechanisms and actionable vulnerabilities

Generation of ACR-368 resistant ovarian cancer cells



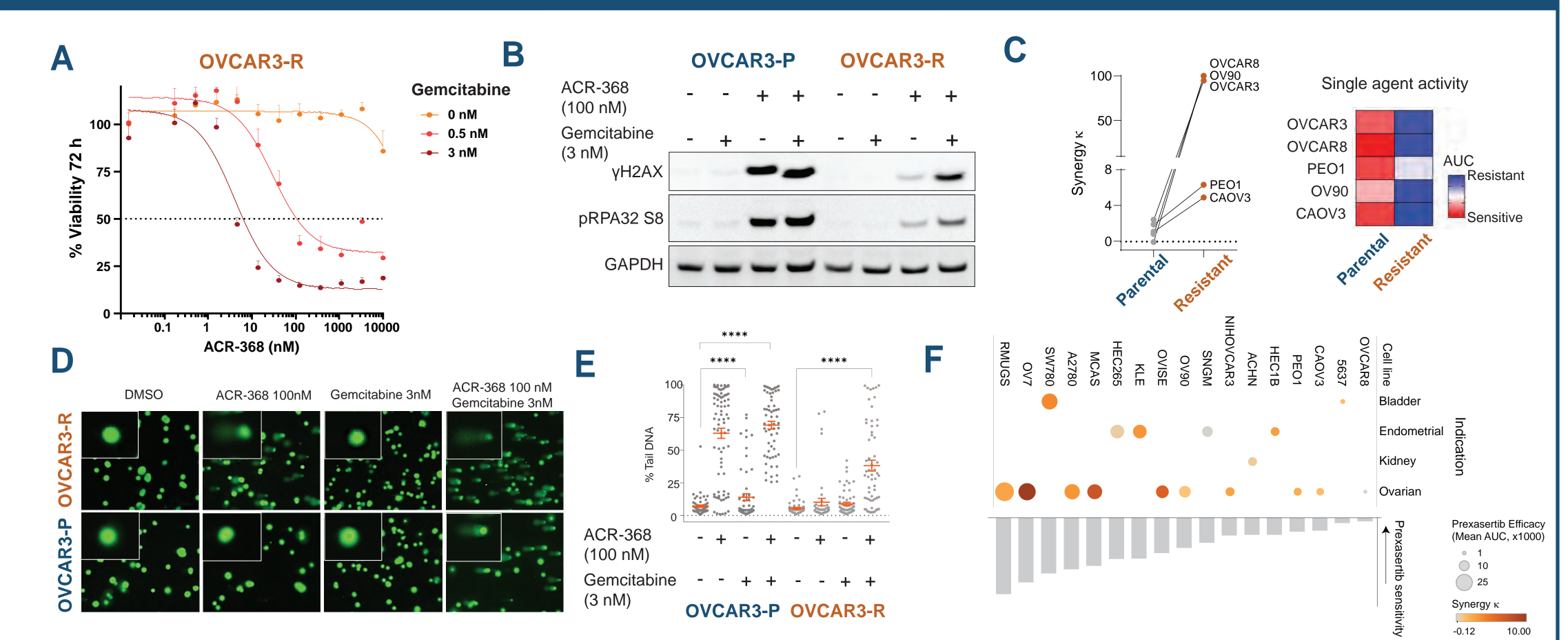
A. OVCAR3 ovarian cancer cells were rendered durably resistant to ACR-368 by culture in continuous presence of 50 nM drug. B. CellTiter Glo assay show drug-induced resistance to ACR-368 in OVCAR3-R cells following 72 hours treatment, compared to OVCAR3 parental cells. C. ACR-368 target engagement was confirmed in both parental and resistant OVCAR3 cells, while induction of DNA damage marker vH2AX was only observed in OVCAR3 parental cells.

AP3 profiling uncovers features linked to ACR-368 resistance



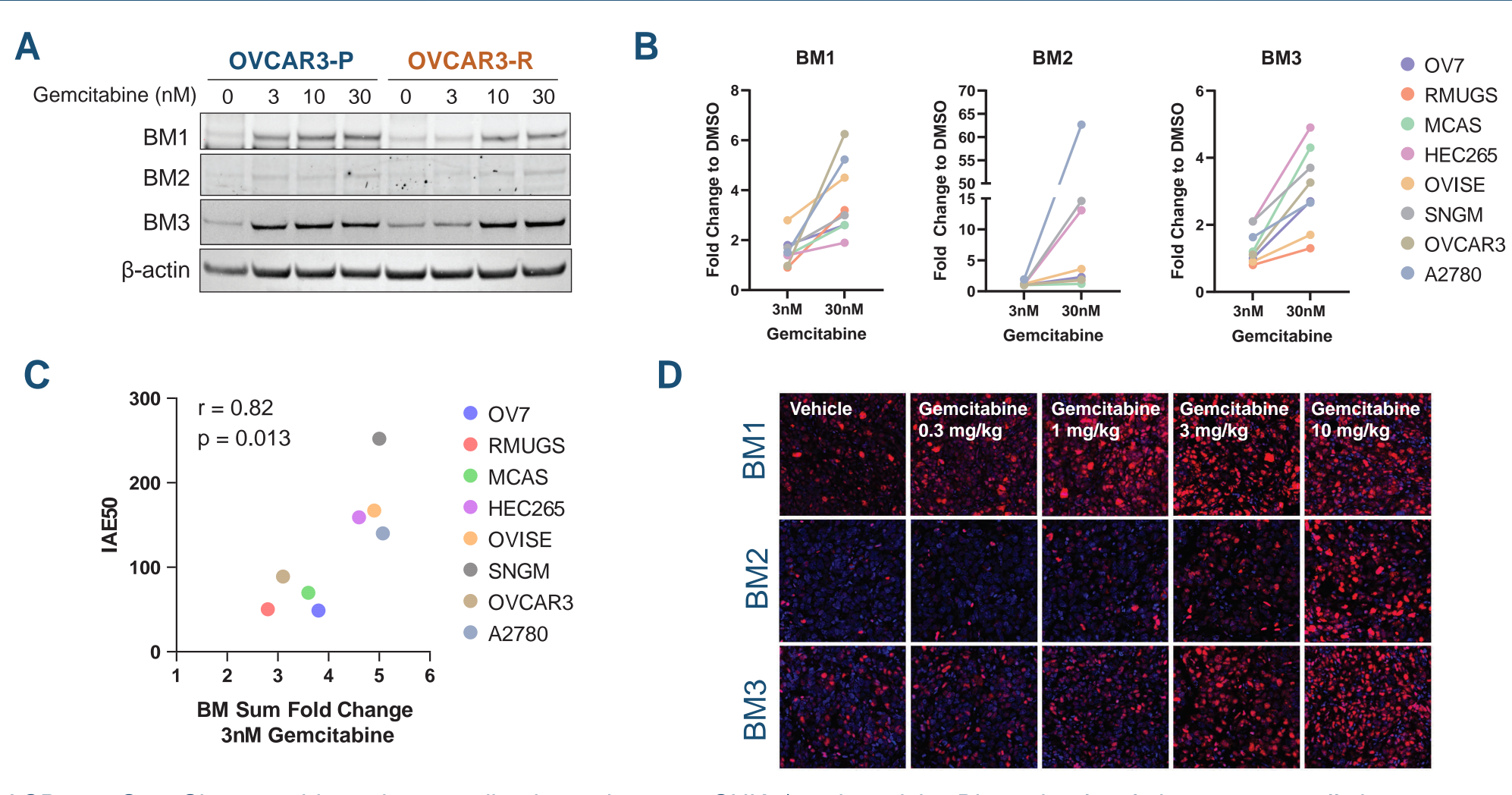
Phosphoproteomic profiling reveals reduced activity around CHK1/2 DDR axis in OVCAR3-R cells. A. Phosphoproteomics workflow. Phosphoproteomes from OVCAR3-P and OVCAR3-R cells treated with ACR-368 were recorded by data independent acquisition mass spectrometry. B. Overview of quantified phosphorylation sites. C. Volcano plots showing ACR-368 regulated phosphosites in relation to DMSO control. D. Top enriched Uniprot keywords associated with regulated phosphoproteins in panel C using Fisher's exact test. E. STRING network visualizations of phosphoproteins underlying terms in panel D. F. Inferred kinase activity networks in ACR-368 treated cells compared to DMSO controls using the Post Translational Modification Set Enrichment Analysis (PTM-SEA) approach. G. Heatmap showing unsupervised hierarchical clustering of proteins with differential abundance in OVCAR3-P and OVCAR3-R cells based on Fisher's exact test - DNA damage repair pathway enriched in OVCAR3-R downregulated cluster.

Low dose Gemcitabine sensitizes ovarian cancer cells to ACR-368



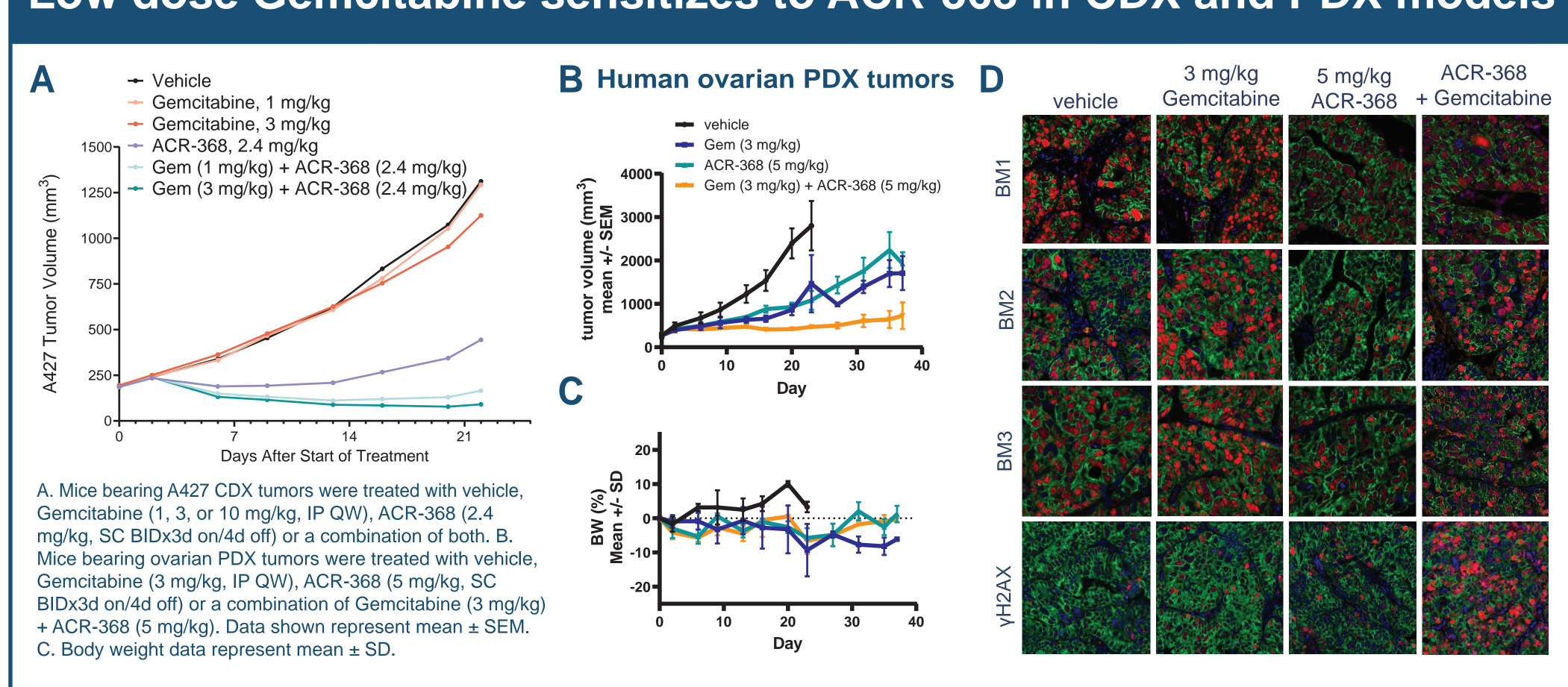
Gemcitabine treatment restores dependency on CHK1/2 activity. A. Gemcitabine sensitize OVCAR3-R cells to ACR-368 in a 72 hours CTG assay. B. Gemcitabine potentiates the effect of ACR-368 and induces markers of DNA damage in OVCAR3-R cells. C. Sensitization to ACR-368 by gemcitabine confirmed in a panel of 5 drug induced ACR-368 resistant cell lines. D. Alkaline comet assay show DNA damage in OVCAR3-R cells treated for 24h with ACR-368 and gemcitabine. E. Quantification of % tail DNA from panel D. F. Sensitization to ACR-368 by gemcitabine validated across a panel of cancer cell lines.

Gemcitabine induces ACR-368 OncoSignature biomarkers



ACR-368 OncoSignature biomarkers predict dependency on CHK1/2 axis activity. Biomarker levels increase as cells become sensitized to ACR-368 by low dose gemcitabine. A-B. Induction of ACR-368 OncoSignature biomarkers evaluated by western blot following 24 hours treatment with gemcitabine in OVCAR3-P and OVCAR3-R cells and across a panel of cancer cell lines. C. Biomarker induction by gemcitabine correlates to combination treatment synergy score in cell line panel. D. Dose-dependent induction of ACR-368 OncoSignature biomarkers in A427 CDX model following 24 hours gemcitabine treatment.

Low dose Gemcitabine sensitizes to ACR-368 in CDX and PDX models



A-C. In vivo efficacy and tolerability of ACR-368 and low dose gemcitabine combination therapy in human non-small cell lung cancer CDX and ovarian PDX models. D. Immunofluorescent staining of ovarian PDX tumors show increased levels of OncoSignature biomarkers following low dose gemcitabine treatment. DNA damage marker vH2AX is strongly induced in tumors following combi-

CONCLUSIONS

- Using AP3 mass spectrometry based pathway mapping of ACR-368 resistance mechanisms, gemcitabine was identified as a rational combination treatment through restoration of stress around the CHK1/2 signal-
- Low dose gemcitabine was confirmed to sensitize cancer cells to ACR-368 across a cell line panel and in CDX and PDX models in vivo
- A concomitant induction of ACR-368 OncoSignature drug response predictive biomarkers was observed following sensitization by gemcitabine treatment in vitro and in vivo
- Efficacy and safety of ultra low dose gemcitabine (10 mg/m²) combined with ACR-368 is currently being evaluated in an exploratory Phase1b/2 clinical study in ACR-368 OncoSignature negative patients
- This study demonstrates the potential of AP3 for unbiased elucidation of actionable drug resistance mechanisms and rapid clinical implementation