ARV-393, a PROteolysis **TArgeting Chimera (PROTAC) BCL6 Degrader, Combined With Biologics or Small-Molecule** Inhibitors Induces Tumor **Regressions in Diffuse Large B-Cell Lymphoma Models**

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Objective

 To assess the activity of the PROTAC B-cell lymphoma 6 (BCL6) degrader, ARV-393, in combination with the standard of care (SOC) first-line chemotherapy regimen for diffuse large B-cell lymphoma (DLBCL), SOC biologics, or small-molecule inhibitors (SMIs) under clinical investigation in DLBCL xenograft models

Key Findings

- ARV-393 in combination with rituximab, cyclophosphamide, hydroxydaunorubicin, vincristine sulfate, and prednisone (R-CHOP), induced significantly greater tumor growth inhibition (TGI) compared with rituximab, CHOP, R-CHOP, or ARV-393 alone, with complete tumor regressions in all mice treated with the combination
- ARV-393 in combination with SOC biologics resulted in superior TGI compared with each agent alone, with complete tumor regressions observed in all mice treated with ARV-393 plus tafasitamab (anti-cluster of differentiation [CD]19) or rituximab (anti-CD20) and an increase in CD20 expression with ARV-393 alone
- ARV-393 in combination with investigational SMIs resulted in superior TGI compared with each agent alone, with tumor regressions observed in all mice treated with the combinations

Conclusions

- ARV-393 demonstrates synergistic antitumor activity, including complete regressions, in combination with SOC agents and select investigational SMIs in high-grade B-cell lymphoma (HGBCL) and aggressive DLBCL models
- These findings support future clinical investigation of ARV-393 in combination with SOC chemotherapy, SOC biologics, and investigational SMIs in patients with **DLBCL**
- Preliminary studies demonstrating that ARV-393 increases CD20 expression provide additional support for the exploration of combinations with CD20-targeted agents and in the context of low or loss of CD20 expression

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Disclosure

All authors are employees and shareholders of Arvinas Operations, Inc. Dan Sherman and Sheryl M Gough additionally hold patents with Arvinas Operations, Inc.

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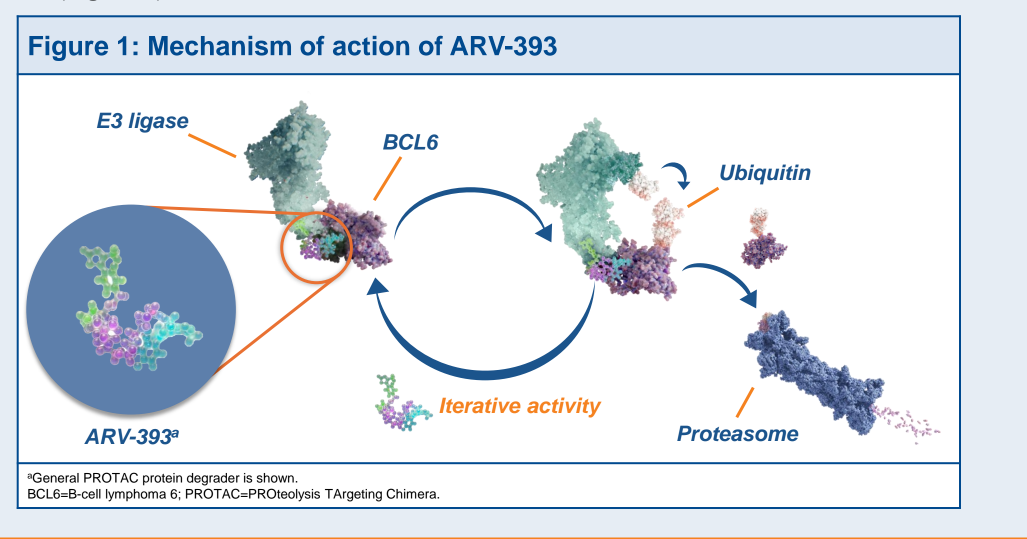
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Background

BCL6 is a preclinically validated oncogenic driver of DLBCL historically considered to be undruggable¹⁻³

Given the heterogeneity and multiple resistance mechanisms of DLBCL and that BCL6 regulates hundreds of genes linked to oncogenesis and resistance,¹ BCL6 degradation has the potential for broad drug combinability

ARV-393, a PROTAC BCL6 degrader, directly binds an E3 ubiquitin ligase and BCL6 to induce the ubiquitination of BCL6 and its subsequent proteasomal degradation (Figure 1)⁴



Results

ARV-393 in Combination With R-CHOP

- The combination of ARV-393 with rituximab, CHOP, or R-CHOP (the first-line SOC therapy for DLBCL) all resulted in tumor regressions; ARV-393 combined with R-CHOP induced complete regressions and had significantly higher TGI compared with rituximab, CHOP, R-CHOP, or ARV-393 alone (Figure 3) - ARV-393 induced complete regressions in 4/10 mice when combined with rituximab, in 6/10 mice when combined with CHOP, and in 10/10 mice when combined with R-CHOP
- Body weights were maintained with monotherapy and combination treatments

ARV-393 in Combination With SOC Biologics

- The combination of ARV-393 with SOC biologics targeting CD19 (tafasitamab), CD79b (polatuzumab vedotin), or CD20 (rituximab) resulted in tumor regressions and demonstrated significantly stronger TGI compared with either agent alone (**Figure 4**)
- ARV-393 combined with tafasitamab induced complete regressions in 10/10 mice (Figure 4A) In contrast, tafasitamab combined with lenalidomide resulted in 55% TGI
- ARV-393 combined with polatuzumab vedotin induced complete regressions in 4/10 mice (Figure 4B)
- ARV-393 combined with rituximab 3 mg/kg or 10 mg/kg induced complete regressions in 9/10 and 9/9 mice, respectively; of note, ARV-393 monotherapy resulted in a significant increase in CD20 expression compared with vehicle (**Figure 4C**)

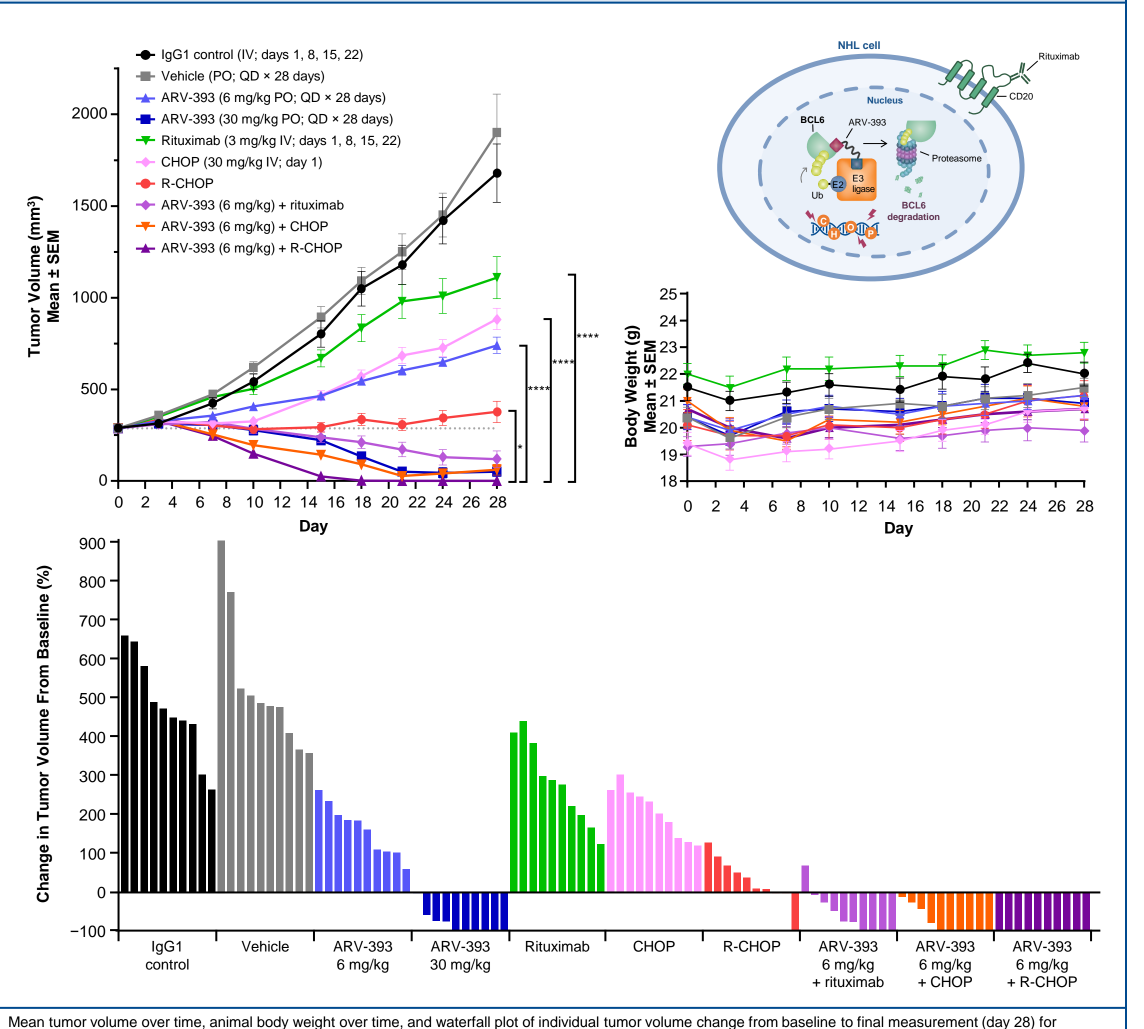
Body weights were maintained with monotherapy and combination treatments

ARV-393 in Combination With SMIs

The combination of ARV-393 with SMIs of BTK (acalabrutinib), BCL2 (venetoclax), or EZH2

- (tazemetostat) demonstrated strong TGI, including tumor regressions in all mice (Figure 5) - ARV-393 combined with acalabrutinib showed significantly stronger TGI than either agent alone (Figure 5A)
- ARV-393 combined with venetoclax demonstrated significantly stronger TGI compared with ARV-393 alone, whereas venetoclax monotherapy resulted in rebound of tumor growth and progressive disease (Figure 5B)
- ARV-393 combined with tazemetostat showed significantly stronger TGI than either ARV-393 or tazemetostat monotherapy (Figure 5C), consistent with literature reports showing that BCL6 and EZH2 play cooperative roles in lymphomagenesis⁷
- In this model, MYC, EZH2, and BCL2 protein levels were increased by 56%, 66%, and 12%, respectively, with ARV-393 alone vs vehicle, but were decreased by 75%, 80%, and 96%, respectively, with ARV-393 plus tazemetostat vs vehicle, demonstrating a synergistic reduction in proteins known to drive lymphoma cell growth
- BCL6 degradation was greater with ARV-393 combined with tazemetostat vs ARV-393 alone (87% vs 65%)
- Body weights were maintained with monotherapy and combination treatments, with dosing holidays implemented in the venetoclax and tazemetostat combinations

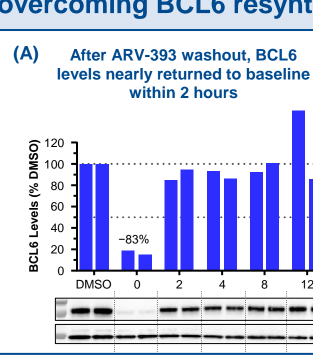
Figure 3: ARV-393 in combination with R-CHOP (SU-DHL-4 model)



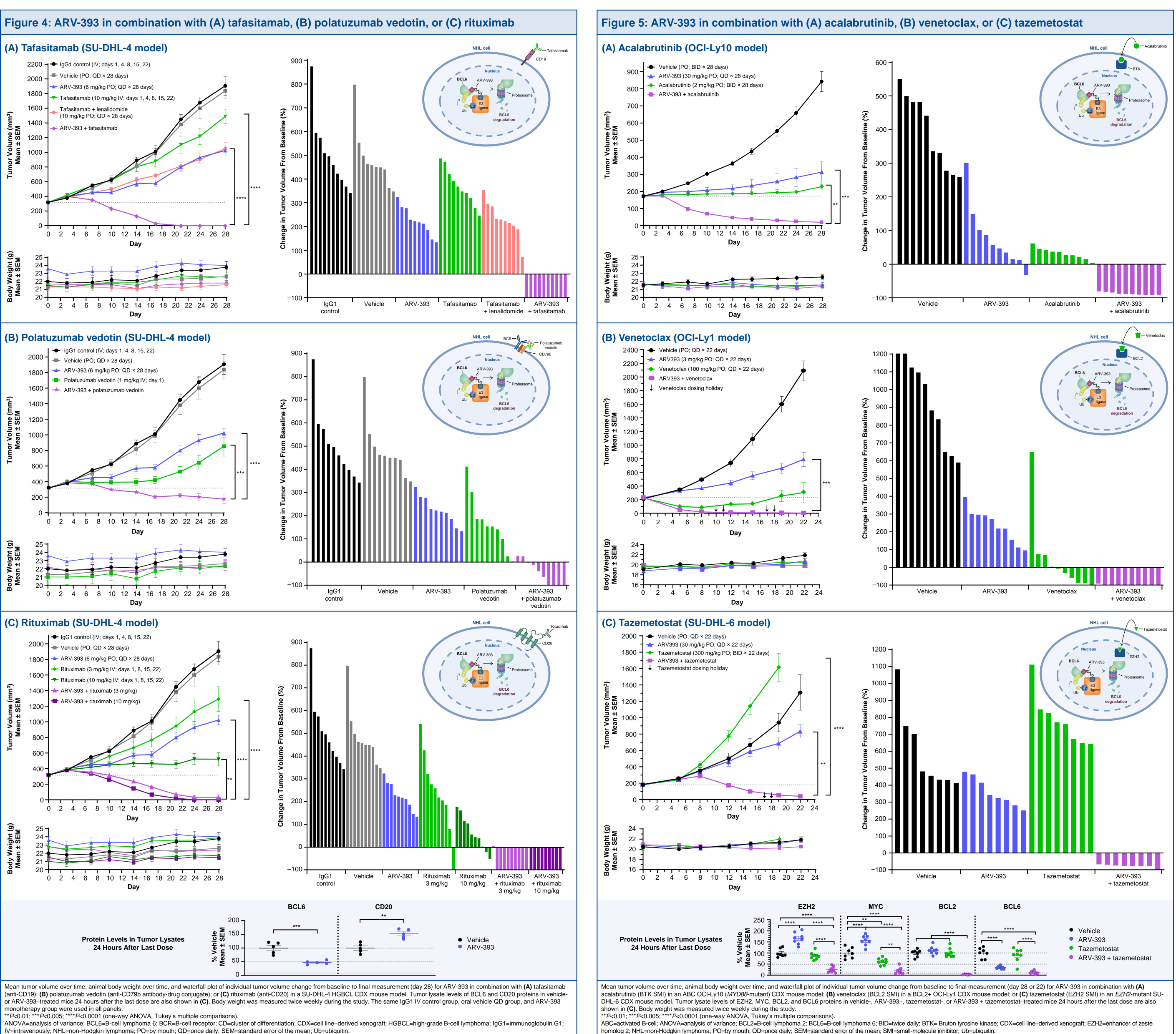
ARV-393 combined with rituximab, CHOP, or R-CHOP in a SU-DHL-4 HGBCL CDX mouse model. Body weight was measured twice weekly during the study. *P<0.05; ****P<0.0001 (one-way ANOVA, Tukey's multiple comparisons) ANOVA=analysis of variance; BCL6=B-cell lymphoma 6; CD=cluster of differentiation; CDX=cell line–derived xenograft; CHOP=cyclophosphamide, hydroxydaunorubicin, incristine sulfate, and prednisone; HGBCL=high-grade B-cell lymphoma; IgG1=immunoglobulin G1; IV=intravenously; NHL=non-Hodgkin lymphoma; PO=by mouth;

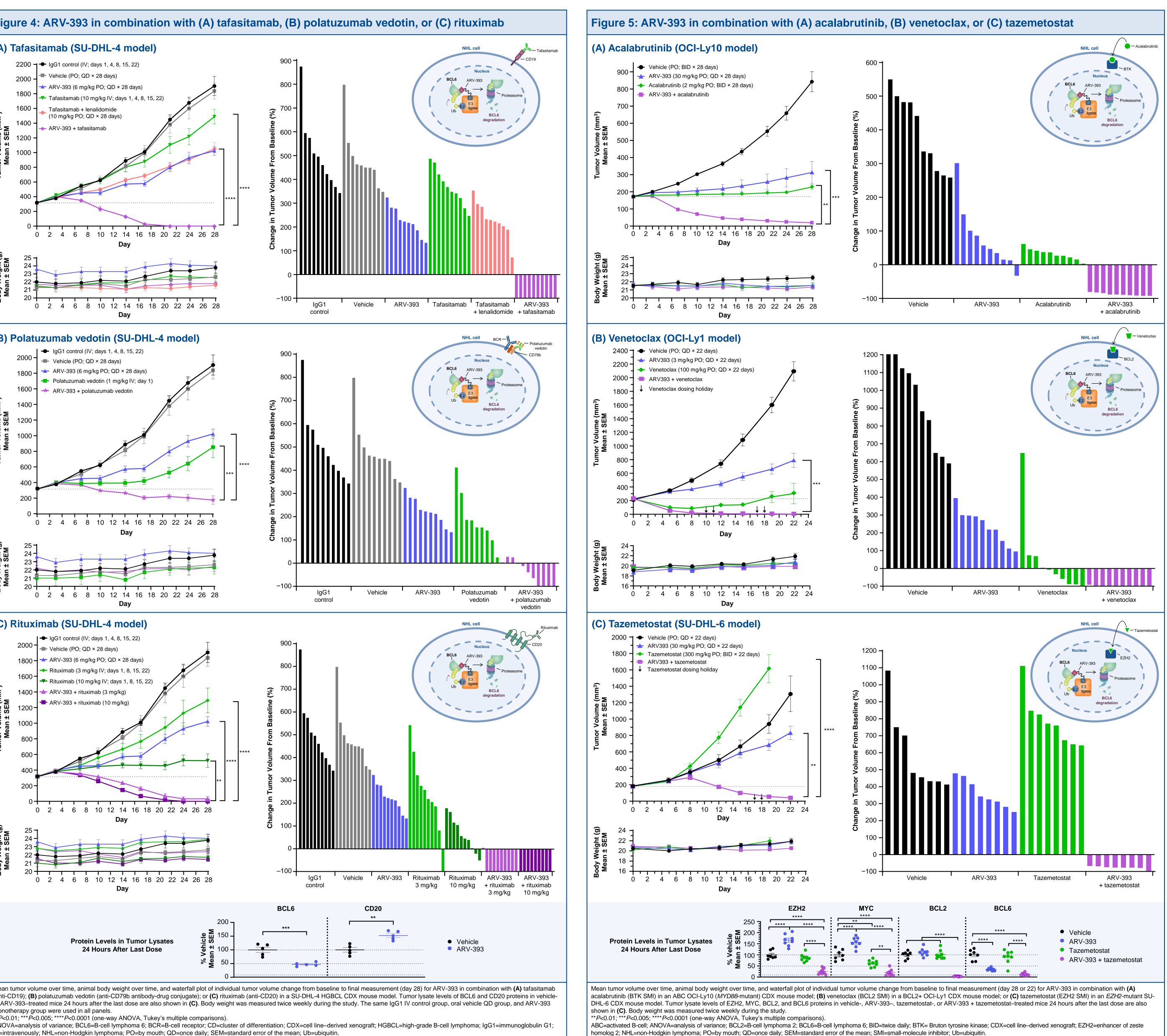
- Hodgkin lymphoma, including DLBCL⁶

Figure 2: ARV-393 potently and rapidly degrades BCL6 in vitro, critical to overcoming BCL6 resynthesis (A) After ARV-393 washout, BCL6 (B) ARV-393 rapidly degrades >90% of BCL6 evels nearly returned to baseline Please scan this within 2 hours QR code to view previously presented ARV-393 DC₅₀ [nM] D_{max} preclinical data.⁵ 🔶 4 hours 24 hours ARV-393 Loq₁₀ [nM (A) BCL6 protein levels quantified by Western blot in the DLBCL OCI-Ly1 cell line, following 4-hour treatment with ARV-393 1.5 nM. Duplicate samples shown following ARV-393 washout and addition of cereblon ligand to block residual ARV-393. (B) BCL6 degradation time-course in OCI-Ly1 cells. BCL6=B-cell lymphoma 6; DC₅₀=half-maximal degradation concentration; DLBCL=diffuse large B-cell lymphoma; D_{max}=maximum level of degradation;



DMSO=dimethyl sulfoxide; GAPDH=glyceraldehyde-3-phosphate dehydrogenase





 ARV-393 rapidly degrades BCL6 in DLBCL cell lines (>90% degradation in 2 hours), and its iterative activity overcomes rapid BCL6 resynthesis (Figure 2); single-agent ARV-393 induced potent TGI, including regressions, in DLBCL patient-derived xenograft models⁵

• ARV-393 monotherapy is being evaluated in a phase 1 trial (NCT06393738) in patients with non-

• Here, we explore the preclinical efficacy of ARV-393 in combination with SOC therapies and SMIs targeting complementary mechanistic pathways in DLBCL

Methods

ARV-393 in Combination With R-CHOP

• A SU-DHL-4 cell line-derived xenograft (CDX) mouse model representing a HGBCL (with MYC, B-cell lymphoma 2 [BCL2], and BCL6 rearrangements) was used to evaluate ARV-393 in combination with rituximab, CHOP, and R-CHOP

- 393 6 mg/kg dose was used for combination studies

ARV-393 in Combination With SOC Biologics

- Using the SU-DHL-4 CDX mouse model, ARV-393 was evaluated in combination with clinically relevant doses of SOC biologic therapies rituximab
- was administered IV on days 1, 8, 15, and 22

combined with tafasitamab

ARV-393 in Combination With SMIs

- 3 mg/kg PO QD was administered alone or in combination with venetoclax (BCL2 SMI)
- One group of mice from each model received the oral vehicle QD

- ARV-393 6 mg/kg or 30 mg/kg was administered orally (PO) once daily (QD) for 28 days; rituximab 3 mg/kg was administered intravenously (IV) on days 1, 8, 15, and 22; CHOP (30:2.475:0.375:0.15 mg/kg) was given IV on day 1 (prednisone was given PO QD on days 1-5); and R-CHOP followed these same dosing methods. The ARV-

- Control groups included mice that received an immunoglobulin G1 (IgG1) IV on days 1, 8, 15, and 22 or mice treated with the oral vehicle QD

- ARV-393 6 mg/kg PO QD was administered alone or in combination with tafasitamab (anti-CD19 biologic), polatuzumab vedotin (anti-CD79b antibody-drug conjugate), or

- Tafasitamab 10 mg/kg was administered IV on days 1, 4, 8, 15, and 22; polatuzumab vedotin 1 mg/kg was administered IV on day 1; and rituximab 3 mg/kg or 10 mg/kg

- Control groups included mice that received IgG1 IV on days 1, 8, 15, and 22; mice treated with the oral vehicle QD; and mice that received lenalidomide 10 mg/kg PO QD

• ARV-393 was evaluated in combination with SMIs in HGBCL or aggressive activated B-cell (ABC) DLBCL CDX models

- ARV-393 30 mg/kg PO QD was administered alone or in combination with acalabrutinib (Bruton tyrosine kinase [BTK] SMI) or tazemetostat (EZH2 SMI); ARV-393

- Acalabrutinib 2 mg/kg PO was administered twice daily (BID) to mice bearing the ABC OCI-Ly10 MYD88-positive CDX, venetoclax 100 mg/kg PO QD to mice bearing the BCL2-positive OCI-Ly1 CDX, and tazemetostat 300 mg/kg PO BID to mice bearing the EZH2-mutant SU-DHL-6 HGBCL CDX