

Editor's Summary

Nanotechnology Sets Sights on Ovarian Tumors

In the world of anticancer research, targeting tumor cells is one challenge; penetrating the cells to deliver therapeutics is another. The combination of specific targeting and efficient delivery is the clinical holy grail, wherein optimization of this approach could lead to highly effective cancer therapy in humans. Ren *et al.* have now developed a nanotechnology platform that allows for just that: targeted intracellular delivery of RNA-based therapeutics to ovarian cancer cells, which halts the oncogenic activity of a potent gene, in this case *ID4*.

In a screen of overexpressed and essential genes in human ovarian cancer, the authors first identified a potential oncogene, *ID4*. They then confirmed *ID4* tumorigenicity and mechanism in vitro in cell lines. After confirming that *ID4* was an oncogene, Ren *et al.* reasoned that "silencing" the gene using small interfering RNA (siRNA) would prevent tumor growth in vivo. The trick was to make sure the siRNA could cross the cell membrane to exert its silencing effects. To accomplish this, the authors designed a tumor-penetrating nanocomplex (TPN) that could not only bind a protein overexpressed on the surface of human cancer cells but also pass through the membrane via a cell-penetrating peptide. Once inside the cells, the TPN could release the siRNA directed against *ID4*. Tumor homing was confirmed in mouse models of human melanoma and ovarian cancer. In mice harboring subcutaneous ovarian tumors, TPN/siRNA decreased *ID4* expression by up to 90% and suppressed tumor growth by 82%. In mice bearing disseminated intra-abdominal tumors, TPN/siRNA allowed 80% of the animals to live 60 or more days. Control treatments did not prevent tumor growth in either study, and the TPN/siRNA therapy did not elicit any immunogenic side effects.

Locked and loaded with siRNA, these TPNs are ready to target and kill cancer cells. The authors envision this to be a platform for credentialing oncogenes and for validating RNA interference in preclinical models before development of therapeutics. However, before moving this TPN/siRNA approach to patients, some additional preclinical optimization is necessary, including pharmacokinetics, testing in human cancer models, and increasing siRNA efficiency at lower doses.

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Targeted Tumor-Penetrating siRNA Nanocomplexes for Credentialing the Ovarian Cancer Oncogene *ID4*

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The comprehensive characterization of a large number of cancer genomes will eventually lead to a compendium of genetic alterations in specific cancers. Unfortunately, the number and complexity of identified alterations complicate endeavors to identify biologically relevant mutations critical for tumor maintenance because many of these targets are not amenable to manipulation by small molecules or antibodies. RNA interference provides a direct way to study putative cancer targets; however, specific delivery of therapeutics to the tumor parenchyma remains an intractable problem. We describe a platform for the discovery and initial validation of cancer targets, composed of a systematic effort to identify amplified and essential genes in human cancer cell lines and tumors partnered with a novel modular delivery technology. We developed a tumor-penetrating nanocomplex (TPN) that comprised small interfering RNA (siRNA) complexed with a tandem tumor-penetrating and membrane-translocating peptide, which enabled the specific delivery of siRNA deep into the tumor parenchyma. We used TPN *in vivo* to evaluate *inhibitor of DNA binding 4 (ID4)* as a novel oncogene. Treatment of ovarian tumor-bearing mice with *ID4*-specific TPN suppressed growth of established tumors and significantly improved survival. These observations not only credential *ID4* as an oncogene in 32% of high-grade ovarian cancers but also provide a framework for the identification, validation, and understanding of potential therapeutic cancer targets.

INTRODUCTION

Genome-scale studies of cancer samples have begun to provide a global depiction of genetic alterations in human cancers, but the complexity and volume of data that emerge from these efforts have made dissecting the underlying biology of cancer difficult, and little is known about the functions of most of the candidates that emerge. For example, in studies of 489 primary high-grade serous ovarian cancer genomes, 1825 genes were identified as targeted by recurrent amplification events (1). Systematic approaches to study the function of genes in cancer cell lines, such as genome-scale pooled short hairpin RNA (shRNA) screens, offer a means to assess the consequences of the genetic alterations found in such genome characterization efforts. We recently used an shRNA-based approach to find genes that are both overexpressed in human primary tumors and essential for the proliferation of ovarian cancer cells (2). This approach identified 54 overexpressed and essential genes in ovarian cancer and 16 genes in non-small cell lung cancer that required further validation *in vivo*.

Furthermore, many of these candidates represent targets that are not amenable to antibody-based therapeutics or traditional small molecule approaches. Thus, if one envisions a discovery pipeline that begins with cancer genomes and ends with novel therapeutics, there is a bottleneck at the point of *in vivo* validation of novel targets.

Achieving silencing in the epithelial cells in the tumor parenchyma is especially critical to study the genetic alterations of interest. RNA interference (RNAi) is a potentially attractive means to silence the expression of candidate genes *in vivo*, particularly for undruggable gene products (3, 4). However, systemic delivery of small interfering RNA (siRNA) to tumors has been challenging, owing to rapid clearance, susceptibility to serum nucleases, and endosomal entrapment of small RNAs, in addition to their inherent inadequate tumor penetration (5). Tumor penetration is also a problem for the delivery of siRNA and shRNA, among other cargos, and is characterized by limited transport into the extravascular tumor tissue beyond the perivascular region (6). This low penetration is thought to arise from the combination of dysfunctional blood vessels that are poorly perfused and a high interstitial pressure, especially in solid tumors, in part due to dysfunctional lymphatics (7). The leakiness of tumor vessels partially counteracts the poor penetration [the so-called enhanced permeability and retention (EPR) effect], but the size dependence and variability of this property can limit its usefulness (7). Desmoplastic stromal barriers can further impede transport of therapeutics through tumors (8, 9).

A new class of tumor-penetrating peptides has been described recently, which home to several types of tumors and leverage a consensus R/KXXR/K C-terminal peptide motif [the C-end rule (CendR)] to stimulate transvascular transport and rapidly deliver therapeutic cargo deep into the tumor parenchyma (10, 11). These peptides are tumor-specific, unlike canonical cell-penetrating peptides (CPPs) that do not display cell- or tissue-type specificity (12), and are able to improve the

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delivery of small molecules, antibodies, and nanoparticles (13). Despite their promise, this class of peptides has not been successfully co-opted for siRNA delivery, in part owing to the additional challenges of delivering oligonucleotides across cell membranes, out of endosomes, and into the cytosol to achieve gene silencing.

Here, we designed an siRNA delivery vehicle that was tumor-penetrating and modular, so it could be easily assembled in a single step to accommodate different payloads to various genes of interest. We envision that such a technology would enable a platform wherein novel targets can be identified by structural and functional genomics and subsequently rapidly credentialed both in vitro and in vivo. Follow-up studies could then identify the mechanism of action underlying the observations and establish (and ultimately prioritize) novel oncogenes as therapeutic targets. To achieve this goal, we have combined a systematic effort to identify genes that are both essential and genetically altered in human cancer cell lines and tumors with the development and deployment of a novel tumor-specific and tissue-penetrating siRNA delivery platform.

RESULTS

Identification of *ID4* as an essential oncogene in human ovarian cancer

To facilitate the identification of genes that are essential in specific cancer types, we initiated Project Achilles, a large-scale effort involving genome-scale pooled shRNA screens in human cancer cell lines (2). Recent efforts to characterize the genomes of primary high-grade serous ovarian cancer have revealed 63 recurrent regions of copy number gain and 50 regions of copy number loss, each containing several genes (1). To identify genes that are both recurrently amplified and essential in ovarian cancers that harbor increased copy number of these genes, we quantified the distribution of shRNA proliferation scores among all shRNAs for each amplified gene (Fig. 1A). We identified 206 cases in which shRNAs targeting the amplified gene were significantly depleted ($P < 0.05$, z score), including known ovarian cancer oncogenes *KRAS*, *AKT1*, *BCL2L1*, and *ERBB3*, and newer candidates for ovarian cancer, including the *inhibitor of DNA binding 4* (*ID4*) transcriptional regulatory protein and the *S-phase kinase-associated protein 2* (*SKP2*) E3-ubiquitin ligase (table S1).

We selected *ID4*, a helix-loop-helix (HLH) transcriptional regulator, for further study because recent genomic analyses have indicated that the chromosomal region containing *ID4* (6p22) is amplified in 32% of high-grade serous ovarian cancers (1) (Fig. 1B). Also, *ID4* is overexpressed in most primary ovarian cancers but not in normal ovary, fallopian tube (FT), and other tissues (Fig. 1C and fig. S1A). In addition, by examining the transcript levels of *ID4* in a large panel of cancer cell lines, we found that *ID4* was frequently overexpressed in most ovarian cancer cell lines and cell lines derived from other cancer lineages, such as endometrial cancer, breast cancer, and glioblastoma (fig. S1B).

After identifying *ID4* as a candidate oncogene in human samples, our next step was to establish preclinical models to credential the oncogenic potential of *ID4*, specifically by investigating the relationship between the amplification, expression, and essentiality of *ID4* in a panel of human cancer cell lines. First, we found that multiple shRNAs that did not alter the expression of *ID1*, *ID2*, and *ID3*, but suppressed *ID4* (fig. S1C), significantly inhibited the proliferation of 9 of 11 ovarian

cancer cell lines and two glioblastoma cell lines tested (range, 53 to 92%) (Fig. 1D). Ovarian cancer cell lines OVCAR-8, OVCAR-4, and CaOV-3, which harbor increased copy number (Fig. 1E) and exhibit overexpression of *ID4* (Fig. 1D), died by apoptosis after *ID4* suppression (fig. S1D). By contrast, seven cell lines that express comparatively lower *ID4* levels showed little inhibition of proliferation after *ID4* suppression (range, 0 to 44%) (Fig. 1D).

We then tested whether *ID4* was an oncogene by investigating its ability to induce cell transformation. Expression of oncogenic *HRAS*^{V12} or coexpression of known RAS effectors renders immortalized human epithelial cells tumorigenic in vivo (14). Therefore, to investigate the role of *ID4* in human ovarian epithelial cells, we similarly created an ovarian surface epithelial cell line [IOSE-M (immortalized ovarian surface epithelial) cells] expressing SV40 large T and small t antigens, hTERT (human telomerase reverse transcriptase), and MEK^{DD} (a constitutively active mutant of a mitogen-activated or extracellular signal-regulated protein kinase kinase) (fig. S1E), and used this cell line to identify genes that promote tumorigenicity. Expression of *ID4* at levels found in ovarian cancer cell lines did not affect the cell cycle progression (fig. S1F) but markedly increased the rate of tumor formation in vivo compared to cells expressing a control vector (Fig. 1F). Addition of *ID4* that harbored mutations in the HLH domain or the addition of *ID1*, *ID2*, or *ID3* each failed to confer significant tumorigenicity (Fig. 1F).

Recent work suggests that most ovarian cancers arise from the FT epithelium rather than the ovarian surface epithelium (15). Hence, we used immortalized FT epithelial cells (16) and assessed whether *ID4* expression also induced cell transformation in these cells. As we found for ovarian epithelial cells, *ID4* induced anchorage-independent growth in FT cells (Fig. 1G). These observations show that *ID4* is amplified, essential, and transforming in ovarian epithelial cells.

ID4-induced transformation through regulation of the *HOXA9* and *CDKN1A* transcriptional programs

To explore the mechanism of action by which *ID4* induces transformation, we performed gene expression profiling in IOSE-M cells that overexpress *ID4* or a control vector. Among the top five gene sets, we found significant enrichment of three gene sets up-regulated by expression of the oncogenic Nucleoporin 98–Homeobox A9 (*NUP98-HOXA9*) fusion protein (17) (Fig. 2A and fig. S2). These three gene sets represent short-term (6 hours) and long-term (3 and 8 days) up-regulation after induction of *NUP98-HOXA9*. *ID4* expression in IOSE-M cells consistently increased the expression of multiple *HOXA* genes, including *HOXA9*, *HOXA7*, and *HOXA3* (Fig. 2B), whereas suppression of *ID4* led to significantly reduced levels of these same genes (Fig. 2C and fig. S3A). To determine whether *HOXA9* was required for *ID4*-induced cell transformation, we introduced three previously validated *HOXA9*-specific shRNAs (18) or control shRNAs targeting green fluorescent protein (*shGFP*) into *ID4*-overexpressing IOSE-M cells (fig. S3B). Suppression of *HOXA9* only slightly (~30%) reduced cell proliferation (fig. S3C) but significantly inhibited *ID4*-induced anchorage-independent growth (Fig. 2D) and reduced the rate of tumor formation in vivo compared to control *shGFP* (Fig. 2E).

Finally, to confirm these observations in human ovarian cancers, we analyzed the expression profiles from primary ovarian tumors by The Cancer Genome Atlas (TCGA) project (1) and found that many genes within the *NUP98-HOXA9* expression signature (TAKEDA_TARGETS OF NUP98-HOXA9 FUSION_10D_DN) were significantly

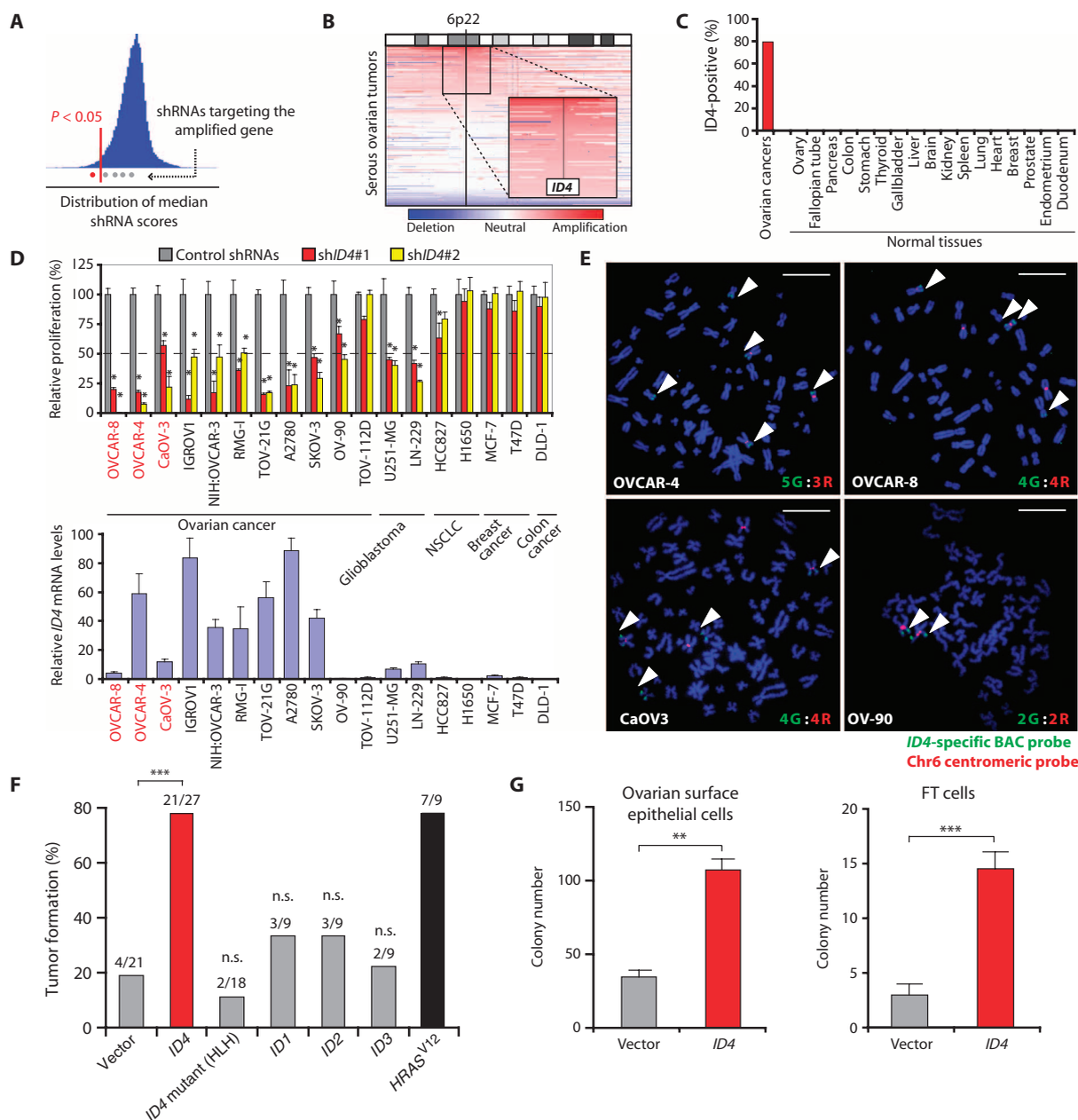


Fig. 1. *ID4* is essential for the proliferation of ovarian cancer cells. **(A)** Median shRNA depletion scores for each amplified gene. Schematic shows the distribution of median shRNA scores (blue bars) in cell lines that harbor copy number gain (\log_2 copy number ratio >0.3) of a given gene. shRNAs targeting the amplified gene (dots) are considered significant if $P < 0.05$ (red line). This analysis was repeated for each of 1825 amplified genes in 63 recurrent regions of genomic amplification identified in primary ovarian tumors. **(B)** Amplification of *ID4* in primary high-grade serous ovarian tumors. SNP array colorgram depicts genomic amplification of *ID4* at chromosome 6p22 region in subsets of primary ovarian tumors, sorted on the basis of the degree of amplification. **(C)** Immunohistochemical analysis of *ID4* performed on sections from tissue microarrays composed of primary human ovarian cancers ($n = 131$) and normal tissues ($n = 85$). **(D)** Effects of

ID4 suppression by two different shRNAs on proliferation of human cancer cell lines (top) and relative levels of *ID4* mRNA (bottom). Data are averages \pm SD ($n = 6$ replicate measurements). 6p22-amplified lines are marked in red. $*P < 0.05$ compared to shGFP (control), Student's *t* test. **(E)** Fluorescence in situ hybridization analysis of *ID4* in ovarian cancer cells. **(F)** Potentiation of tumorigenicity by *ID4* overexpression. IOSE-M cells expressing the indicated constructs were implanted subcutaneously into immunodeficient mice. The number of tumors formed/number of injections is indicated. *HRAS*^{V12}-expressing IOSE-M cells were used as a positive control. $***P < 0.001$, Fisher's exact test; n.s., not significant. **(G)** *ID4* promotes anchorage-independent growth of IOSE-M (ovarian epithelial) and FTSEC-M (FT) cells. Data are averages \pm SD ($n = 6$). $**P < 0.01$, $***P < 0.001$ compared to respective vector control, Student's *t* test.

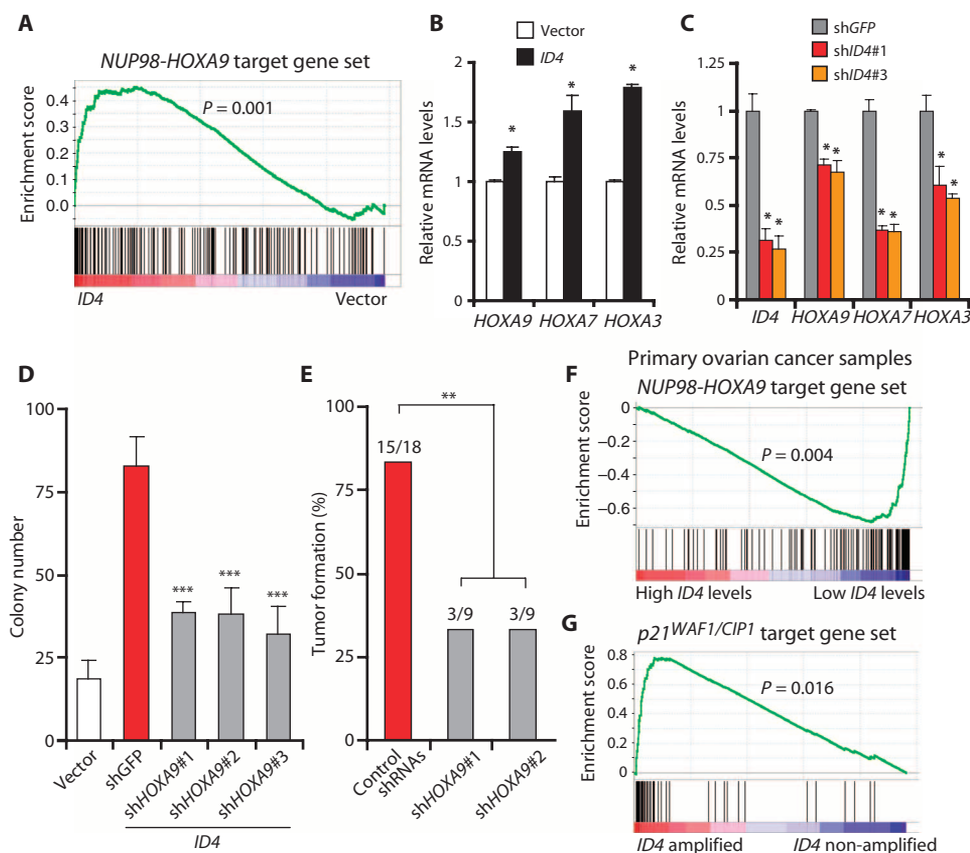


Fig. 2. *ID4* induces tumorigenicity that depends on *HOXA9*. (A) *ID4* induces *HOXA9* gene activity. Gene expression profiling and gene set enrichment analysis (GSEA) were performed on IOSE-M cells overexpressing *ID4* or a control vector. All genes were ranked on the basis of the differential expression between cells expressing *ID4* or a control vector. Black bars at the bottom of the figure indicate the location of genes in an *NUP98-HOXA9* up-regulated gene set (TAKEDA_TARGETS_of_NUP98_HOXA9_FUSION_3D_UP) (17) within the ranked list, and the green curve indicates the running enrichment score for the gene set. (B) Quantitative reverse transcription–polymerase chain reaction (RT-PCR) analysis of *HOXA9*, *HOXA7*, and *HOXA3* mRNA in IOSE-M cells overexpressing *ID4* or a control empty vector. Data are averages \pm SD ($n = 6$ replicate measurements). (C) Quantitative RT-PCR analysis of *HOXA9*, *HOXA7*, and *HOXA3* mRNA in OVCAR-8 cells 3 days after infection with a control shGFP or two shRNAs targeting *ID4*. Data are averages \pm SD ($n = 6$ replicate measurements). (D) Anchorage-independent growth of *ID4*-overexpressing IOSE-M cells in response to sh*HOXA9*. Data are averages \pm SD ($n = 6$). *** $P < 0.0001$, Student's t test. (E) Effect of *HOXA9* suppression on *ID4*-induced tumorigenicity in immunodeficient mice. *ID4*-overexpressing IOSE-M cells expressing indicated shRNAs were subcutaneously implanted into immunodeficient mice. The number of tumors formed/number of injections is indicated. ** $P < 0.01$, Fisher's exact test. (F) Expression data from primary ovarian tumors with low *ID4* expression levels ($n = 44$ samples) were compared to samples with high *ID4* expression ($n = 45$). Thresholds were 1 SD below and above the mean expression of all the samples. At the bottom of the enrichment plots, black bars indicate the location of genes in an *NUP98-HOXA9*-down-regulated gene set (TAKEDA_TARGETS_OF_NUP98_HOXA9_FUSION_10D_DN) (17). (G) *ID4* amplification in ovarian tumors correlated with decreased p21^{WAF1/CIP1} activity. Expression profiling of primary ovarian tumors with matched copy number data was used to perform GSEA on amplified *ID4* (\log_2 copy number ratio >0.3) and nonamplified *ID4* samples (\log_2 copy number ratio <0). All genes were ranked by their differential expression (signal to noise) between 81 nonamplified and 109 amplified *ID4* primary tumors. Black bars at the bottom of the figure indicate the location of genes in the p21^{WAF1/CIP1} target gene set (P21_P53_ANY_DN).

down-regulated in tumors that express low levels of *ID4* (Fig. 2F). The gene set was originally defined 10 days after expression of *NUP98-HOXA9*. Human tumors that harbored *ID4* amplifications showed significantly increased expression of a gene set that was down-regulated by p21^{WAF1/CIP1} or TP53 tumor suppressor (P21_P53_ANY_DN) (19),

terminus to facilitate interactions with membrane lipids (Fig. 3B and fig. S4A). We first confirmed that peptides from the library bound siRNA electrostatically to form stable nanocomplexes in water (diameter, ~ 80 nm), phosphate-buffered saline (PBS, diameter ~ 200 to 400 nm), and serum (diameter, ~ 200 to 400 nm) (fig. S4, B and C). Most peptides

indicating a suppressed p21^{WAF1/CIP1} transcriptional activity in *ID4*-amplified tumors (Fig. 2G). These observations demonstrate that expression or amplification of *ID4* regulates the *HOXA9* and *CDKN1A* (cyclin-dependent kinase inhibitor 1A) transcriptional programs to disrupt the normal regulation of proliferation and differentiation.

Design of tumor-penetrating nanocomplexes

To assess whether *ID4* functions as a therapeutic target for ovarian cancer, we designed an siRNA delivery platform that permits access to specific cancer cells. To this end, we synthesized a library of tandem peptides capable of tumor homing and penetration (with a fixed cyclic domain) and siRNA delivery across cell membranes (variable linear domains for electrostatic binding) (Fig. 3A and table S2). We selected the cyclic nonapeptide LyP-1 (CGNKRTRGC) as the tumor-penetrating domain for all tandem peptides because it binds p32 [p33/gC1q receptor/hyaluronan binding protein 1 (HABP1)], a mitochondrial protein whose expression is elevated on the surface of stressed tumor and tumor-associated cells in a wide range of tumor types (20). Once bound, LyP-1 is proteolytically processed by endogenous proteases to reveal a cryptic CendR motif that activates tissue penetration of associated payloads and results in higher therapeutic accumulation (11, 21).

We postulated that a tandem peptide consisting of an N-terminal CPP with a C-terminal LyP-1 separated by a four-glycine linker domain would retain membrane translocation and tissue-penetrating properties, condense siRNA into nanoparticles, and selectively target tumor cells expressing p32 at the surface. We created a library of 18 linear membrane translocation domains that contained both polycationic peptide sequences [oligoarginines of varying net charge, TAT(48–60), and the herpes simplex virus type 1 tegument protein VP22] and amphipathic sequences [penetratin (PEN) and transportan (TP)]. Inspired by nonenveloped RNA viruses (22), a myristoyl group was added to the amino

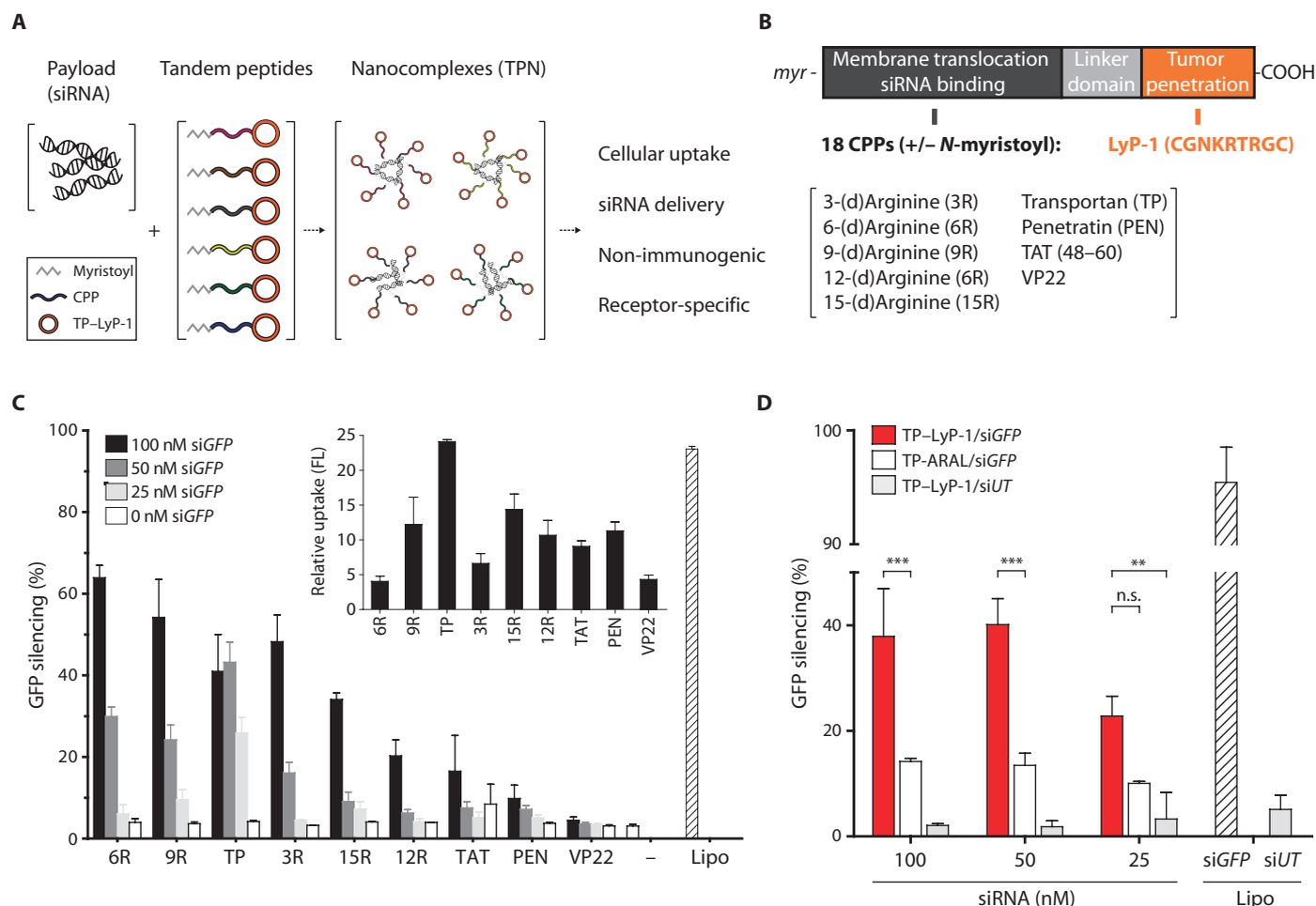


Fig. 3. Characterization and activity of TPNs. **(A)** Schematic of the tandem peptide screen. siRNA was noncovalently bound to a library of 18 candidate tandem peptides bearing a fixed N-terminal cyclic tumor penetration domain and variable C-terminal linear membrane translocation domains. The resulting nanocomplexes were assayed and selected for their cellular uptake, siRNA delivery, lack of immunogenicity, and receptor specificity in human cancer cell lines in vitro or in mouse models of human ovarian cancer. **(B)** The tandem peptide construct and various membrane translocation domains tested. **(C)** Gene silencing activity of tandem peptide–siGFP nanocomplexes in HeLa cells stably expressing GFP. Percent GFP silencing was calculated on the basis of the geometric mean of GFP fluorescence intensity of the whole population relative to cells treated with media only. The Lipofectamine control (Lipo) contained 100 nM GFP–siRNA. Data are averages ± SD (*n* = 4 independent experiments). Inset: HeLa cell uptake of fluorescently labeled nanocomplexes was assessed by flow cytometry. Data are averages ± SD (*n* = 4 independent experiments). **(D)** Receptor specificity of nanocomplex-mediated GFP silencing. Data are averages ± SD (*n* = 6 independent experiments). ****P* < 0.01, *****P* < 0.001, one-way analysis of variance (ANOVA).

were noncytotoxic, with the exception of highly cationic peptides 12R and 15R (fig. S4D).

To evaluate nanocomplex-mediated gene silencing, we treated HeLa cells expressing destabilized GFP with siRNA targeting GFP (siGFP) bound to either tandem peptides or Lipofectamine. The top three carriers—6R, 9R, and TP—silenced GFP expression 40 to 60% (Fig. 3C). Flow cytometry analysis indicated that gene suppression was dose-dependent and heterogeneous across the cell population, with 50 to 70% of cells exhibiting near-complete gene suppression (fig. S4E), with variation likely owing to heterogeneity in p32 expression. GFP suppression can underreport the magnitude of silencing endogenous gene targets, such as amplified oncogenes (23); thus, we used this assay to compare peptides but not as a quantitative measure of efficiency of RNAi of oncogenes. We chose nanocomplexes consisting of siRNA bound to TP–LyP-1 for further analysis [hereinafter referred to as the tumor-

penetrating nanocomplex (TPN)] because it had the highest efficacy in delivering siRNA to cells, was potent relative to other tandem peptides (>25% gene suppression at 25 nM siRNA), and has a known secondary structure (24), in contrast to poly-Arg peptides. Next, we examined the specificity of receptor targeting and gene suppression. Uptake of LyP-1–targeted nanocomplexes was significantly higher than control nanocomplexes bearing ARAL (ARALPSQRSR), a peptide that has the same net charge as LyP-1 (fig. S4F). LyP-1 targeting was specific, as shown by competition assays with free LyP-1 (fig. S4G) and in cells lacking the p32 receptor (fig. S4H). We did not detect statistically significant alterations in GFP expression in HeLa cells exposed to untargeted control nanocomplexes (UCNs) carrying siGFP (TP-ARAL/siGFP) or targeted TP–LyP-1 carrying a scrambled siRNA sequence (TP–LyP-1/siUT) (Fig. 3D). In contrast, when we used TP–LyP-1 carrying siGFP, we observed more than 40% suppression of GFP expression.

OVCAR-4 and OVCAR-8 human ovarian cancer cell lines have elevated surface p32 expression (fig. S5); therefore, these cell lines were compatible for testing TPN-mediated delivery of *ID4*-specific siRNA (*siID4*). When *ID4* was suppressed in these cell lines in vitro using TPN (Fig. 4A), we observed a decrease in cell viability (Fig. 4B) and an increased rate of apoptosis (Fig. 4C and fig. S5). We therefore selected the OVCAR-4 and OVCAR-8 ovarian cancer cell lines for our preclinical evaluation of *ID4* in vivo.

TPN homing and penetration in vivo in mice

For in vivo applications, an ideal carrier should extend the half-life of siRNA in serum by protecting it from nuclease degradation, yet readily dissociate from siRNA once in the cytosol of target cells. We used a gel-shift assay to determine an optimal siRNA-to-peptide complexation molar ratio of 1:20, at which free siRNAs were encapsulated into TPN, likely owing to charge-based interactions between the peptide and the siRNA backbone (Fig. 5A). We found that TPN extended the stability of siRNA to >12 hours in murine serum (Fig. 5A), and the size of TPN remained stable in the presence of serum over the course of 10 hours (Fig. 5B). Furthermore, we observed unpackaging of TPN upon exposure to endolysosomal pH (5.3 to 6.5) (Fig. 4D).

We assessed whether TPN enabled receptor-specific, tumor-penetrating siRNA delivery to established tumors (Fig. 5C) by examining their in vivo homing behavior in mice bearing subcutaneous human melanoma tumors (MDA-MB-435 xenografts). After intravenous administration, most siRNAs were cleared renally, as indicated by bladder accumulation of siRNA over 8 hours (fig. S6A). However, we also observed that TPN was cleared more slowly than naked siRNA (Fig. 5D) and distributed to tumor tissues within 30 min (fig. S6B). We noted similar pharmacokinetics for intravenous and intraperitoneal injections of TPN, suggesting the potential for multiple routes of administration (Fig. 5D and fig. S6A).

Biodistribution studies showed blood clearance through liver and spleen accumulation, likely owing to reticuloendothelial uptake. TPN also distributed to the lung (fig. S6, C and D), a common uptake site for passively targeted nanoparticles, owing to interactions with the first capillary bed after intravenous injection (25). After clearance from circulation (4 hours), we found a more than threefold increase in the tumor fluorescence of TPN in MDA-MB-435 xenografts compared to that of UCNs (Fig. 5E). Similar results in tumor targeting were obtained in mice bearing OVCAR-8 ovarian tumor xenografts (fig. S6E).

Although bulk tumor accumulation provides a global measure of targeting, it does not provide spatial information on the extravascular and tumor interstitial availability of TPN, which ultimately determines their efficacy. We thus performed histological characterization of TPN penetration in xenografted subcutaneous OVCAR-8 tumors at various time points and found that these TPN exhibited an initial intravascular distribution and subsequently extravasated into the tumor interstitium (Fig. 5F). The fraction of TPNs that were sequestered beyond the intravascular space was significantly higher than that of UCNs (Fig. 5F). Furthermore, TPNs accumulated in the tumor parenchyma to a degree similar to that of nanocomplexes bearing a different tumor-penetrating motif, immobilized arginine-glycine-aspartate (iRGD) (CRGDKGPDC) (10). Nonpenetrating, targeted nanocomplexes (TP-RGD4C) (26) or the commercial reagent Lipofectamine failed to show a similar distribution (Fig. 5G). We also found that TPN also colocalized with p32-positive tumor cells, as expected, but failed to detect UCNs or naked siRNAs in tumors (fig. S6F). This pattern of distribution is

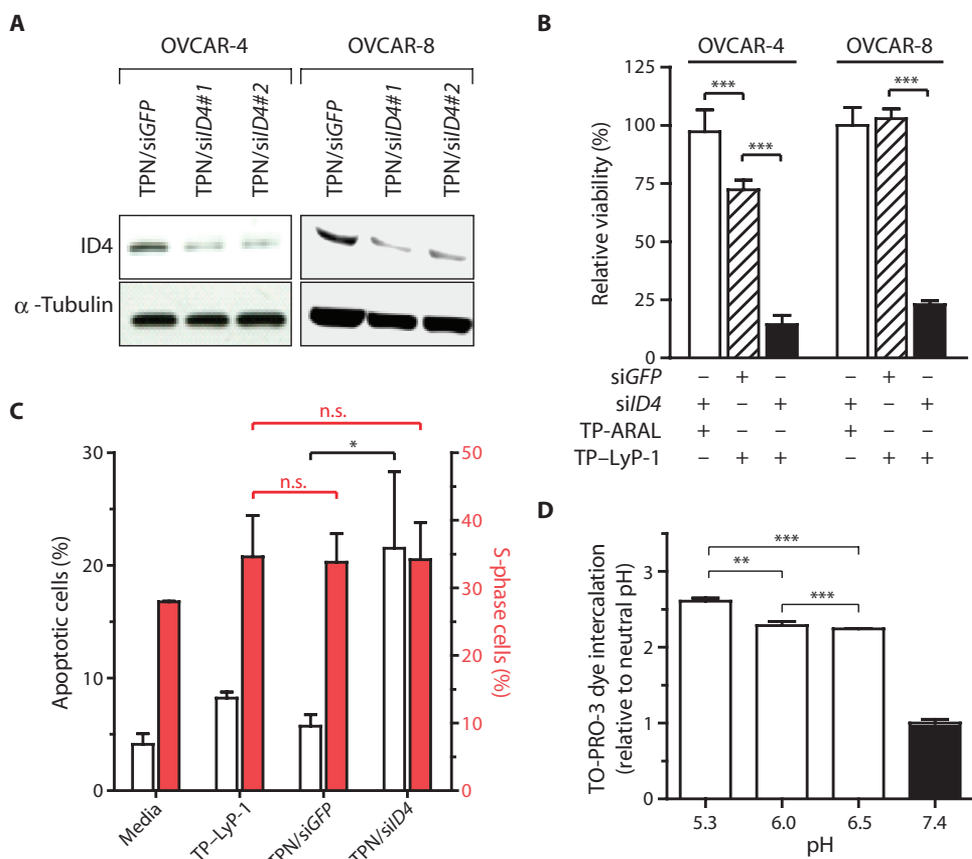


Fig. 4. TPN-mediated suppression of *ID4* in p32-expressing ovarian cancer cells. **(A)** *ID4* suppression by TPN-mediated siRNA delivery in vitro. Immunoblot of *ID4* in two p32-amplified ovarian cancer cell lines, OVCAR-4 and OVCAR-8, which were treated with TPN containing one of two *ID4*-specific siRNAs or a control siRNA targeting GFP (*siGFP*). α -Tubulin was used as a loading control. **(B)** Effects of *ID4* suppression on cell proliferation. Data are averages \pm SD ($n = 4$ independent experiments). *** $P < 0.001$, one-way ANOVA. **(C)** Effects of *ID4* suppression in OVCAR-8 cells. The percentages of apoptotic and S-phase cells were calculated. Data are averages \pm SD ($n = 3$ independent experiments). * $P < 0.05$, one-way ANOVA. **(D)** Intercalation of TO-PRO-3, a nucleic acid-binding dye, into siRNA in the presence of TPN at various pH. Fluorescence was detected at 640/680 nm (excitation/emission). Data are averages \pm SD ($n = 4$ independent experiments). ** $P < 0.01$, *** $P < 0.001$, one-way ANOVA and Tukey post hoc tests.

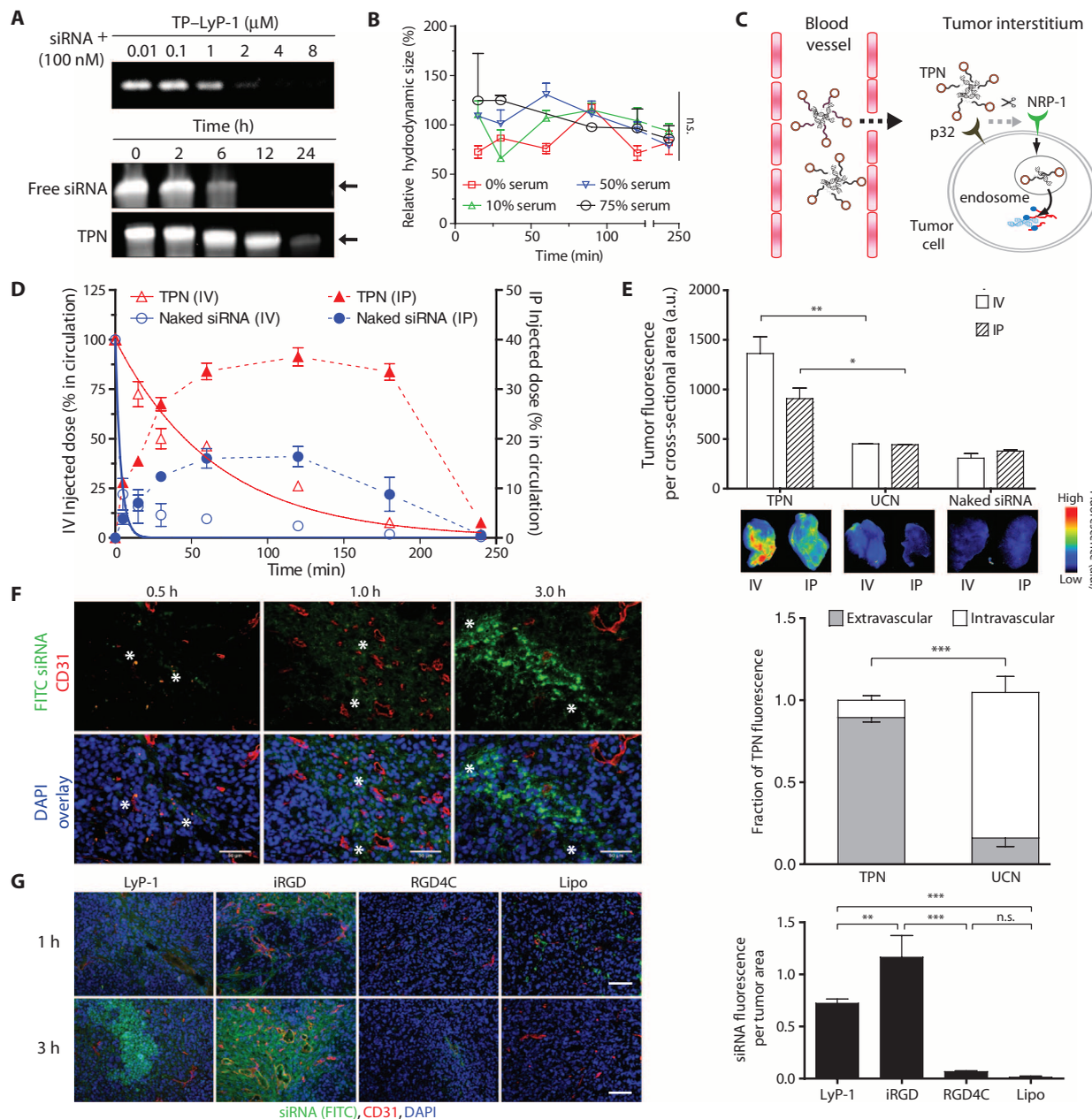


Fig. 5. TPN homing in vivo. **(A)** Serum stability of TPN. (Top) Agarose gel analysis of siRNA complexed to TP-LyP-1 at varying molar ratios. (Bottom) Gel electrophoresis of free siRNA and TPN in mouse serum at 37°C at the indicated times. Arrow indicates expected position of intact siRNA (14 kD). **(B)** Dynamic light scattering measurements of TPN in varying concentrations of mouse serum at 37°C over time. The hydrodynamic diameter of TPN in each serum concentration over time is normalized to its size in PBS (0% serum) at time 0 min. Data are averages ± SD (*n* = 6 independent measurements), one-way ANOVA. **(C)** Schematic of TPN penetration and targeted delivery of siRNA into the cytosol of cancer cells. **(D)** In vivo circulation of siRNA compared with naked siRNA against GFP upon intravenous (IV) or intraperitoneal (IP) administration. Fluorescence of siRNA was detected in the blood drawn retro-orbitally. Data are averages ± SD (*n* = 3). **(E)** Quantification of siRNA fluorescence and corresponding fluorescence images of MDA-MB-435 whole-tumor explants harvested 4 hours after injection of TPN, UCN, or naked siRNA, either intravenously or intra-

peritoneally. Data are averages ± SD (*n* = 6). **P* < 0.05; ***P* < 0.01, one-way ANOVA. **(F)** Histological analysis of time-dependent homing of TPN carrying fluorescein isothiocyanate (FITC)-labeled siRNA (asterisks) in relation to blood vessels in mice bearing human OVCAR-8 tumor xenografts. Nuclei were stained with 4',6-diamidino-2-phenylindole (DAPI). Tumor vasculature is CD31⁺. Scale bars, 50 μm. Right: Extravascular and intravascular fractions of TPN were quantified from the fluorescence images. Data are averages ± SD, from representative sections of six independent tumors. ****P* < 0.001 by two-tailed Student's *t* test. **(G)** Tumor parenchyma penetration by TPN with LyP-1 and iRGD homing domains, nonpenetrating nanocomplex targeted by RGD4C peptide, and Lipofectamine. Nanocomplexes were injected intravenously, and OVCAR-8 tumors were stained 1 or 3 hours later (*n* = 6 per formulation). Scale bars, 50 μm. Right: Tumor fluorescence was quantified. Data are averages ± SD from six randomly selected views per condition. ***P* < 0.01, ****P* < 0.001, one-way ANOVA with Tukey post hoc tests.

similar to previously reported LyP-1 peptide (21) and LyP-1-targeted nanoparticle homing (27).

Loss of tumor maintenance after TPN-mediated suppression of *ID4* in vivo

We next tested whether *ID4* was essential for tumor maintenance through TPN-mediated delivery of *siID4* in established human ovarian cancer xenografts in vivo. Mice harboring subcutaneous OVCAR-4 tumors were injected intravenously or intraperitoneally with TPN/*siID4* every 3 days for 25 days. Control animals with tumors were injected intravenously with saline, TP-LyP-1 carrier without siRNA, or TPN/*siGFP*. Repeated, systemic administration of the TPN/*siID4* resulted in 80 to 90% decrease in *ID4* mRNA relative to treatment with TPN/*siGFP* (Fig. 6A), with a corresponding suppression of tumor growth by 82% in intravenously injected animals and 87% in intraperitoneally injected animals (Fig. 6, B and D). Tumor growth was static in *siID4*-treated cohorts for 30 days after the termination of TPN treatment (Fig. 6B). In contrast, TPN/*siGFP* or TP-LyP-1 carrier had no effect on *ID4* and did not prevent further tumor growth. We also found that both methods of TPN/*siID4* injection induced a significant increase in *CDKN1A* (p21^{WAF1/CIP1}) transcript levels (Fig. 6C), a known target of *ID4* in mediating cell cycle arrest (28). Residual TPN/*siID4*-treated tumors exhibited significant apoptosis (Fig. 6E).

We then tested whether TPN/*siID4* could induce therapeutic activity in mice harboring disseminated tumors. We established orthotopic OVCAR-8 tumors, confirmed intra-abdominal dissemination 3 weeks after implantation, and then initiated intraperitoneal injection of TPN every 3 days for 30 days. The tumor burden in mice that received TPN/*siID4* remained low compared to controls (Fig. 7A and fig. S7A), and 80% of the recipients of TPN/*siID4* (intraperitoneal) survived more than 60 days, even after the nano-complex treatments ceased at day 50 (Fig. 7B). After 40 days, disseminated tumors and hemorrhagic ascites were present in control cohorts (fig. S7B), but there was no visible tumor lesions in four of five TPN/*siID4*-treated mice upon necropsy, indicating tumor regression. Histological analysis of the single visible remnant tumor in an *siID4*-treated animal revealed significant reduction in *ID4* levels relative to tumors from the three control cohorts and increased apoptosis in the tumor parenchyma (Fig. 7C and fig. S7C).

The versatility of TPN for direct target comparison was demonstrated by treating mice bearing OVCAR-8 tumors with TPN carrying siRNA against *Claudin-3* (*CLDN3*),

a recently reported potential ovarian siRNA target (29). TPN/*siCLDN3* led to initial reduction of tumor growth compared to controls treated with saline or TP-LyP-1 carrier alone but was unable to achieve sustained growth suppression compared to TPN/*siID4* treatments (fig. S7D).

Side effects of TPN administration in vivo

In some cases, apparent therapeutic effects of RNAi may be confounded by non-sequence-specific innate immune responses (30). Therefore, we independently measured serum levels of interferon- α , tumor necrosis factor- α , and interleukin-6 in immunocompetent mice after administering various TPN-based siRNA formulations and failed to observe induction of these cytokines in animals that received TPN/*siID4*. By comparison, mice that received an immunostimulatory siRNA, *si β gal-728* (30), complexed to either Lipofectamine or TP-LyP-1 were positive controls for nonspecific immunostimulation (fig. S8A). Moreover, we did not observe deleterious effects on animal weight in two ovarian tumor models (fig. S8B) or evidence of macro-

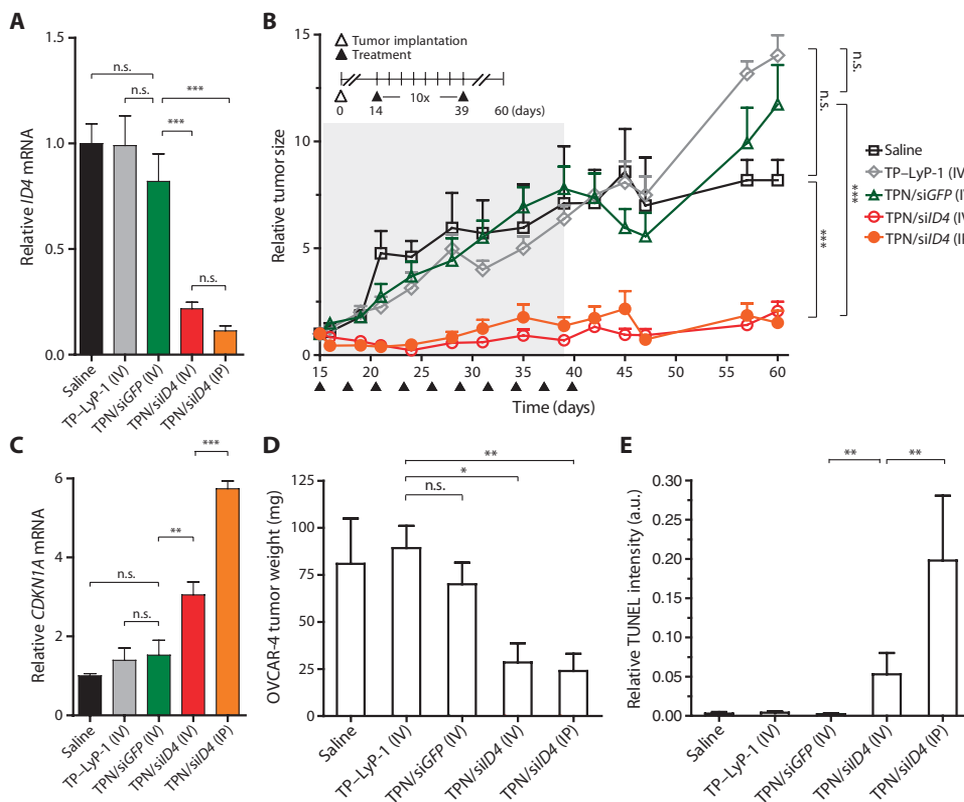


Fig. 6. Subcutaneous tumor treatment with TPN/*siID4*. **(A and B)** Efficacy of TPN-mediated delivery of *siID4* in vivo. *ID4* mRNA (A) and burden of subcutaneous OVCAR-4 xenografts (B) after treatments with the indicated formulations every 3 days for 25 days (arrowheads). Control cohorts received either TP-LyP-1 without siRNA or TPN/*siGFP*. Inset shows the experimental timeline. Treatment period is shaded in gray. Data are averages \pm SD ($n = 8$ to 10 tumors per group). *** $P < 0.001$, one-way ANOVA. **(C)** *CDKN1A* mRNA levels from tumors at day 60, relative to saline control. Data are averages \pm SD. *** $P < 0.01$; **** $P < 0.001$, one-way ANOVA. **(D)** Weight of OVCAR-4 tumors at day 60. Data are averages \pm SD ($n = 5$ to 10 tumors per cohort). * $P < 0.05$, ** $P < 0.01$, one-way ANOVA. **(E)** Quantification of TUNEL (terminal deoxynucleotidyl transferase-mediated deoxyuridine triphosphate nick end labeling) staining intensities from 6 to 10 randomly selected OVCAR-4 tumor sections ($n = 5$ per treatment group) after 30 days of TPN treatment. Data are averages \pm SD. ** $P < 0.01$, one-way ANOVA.

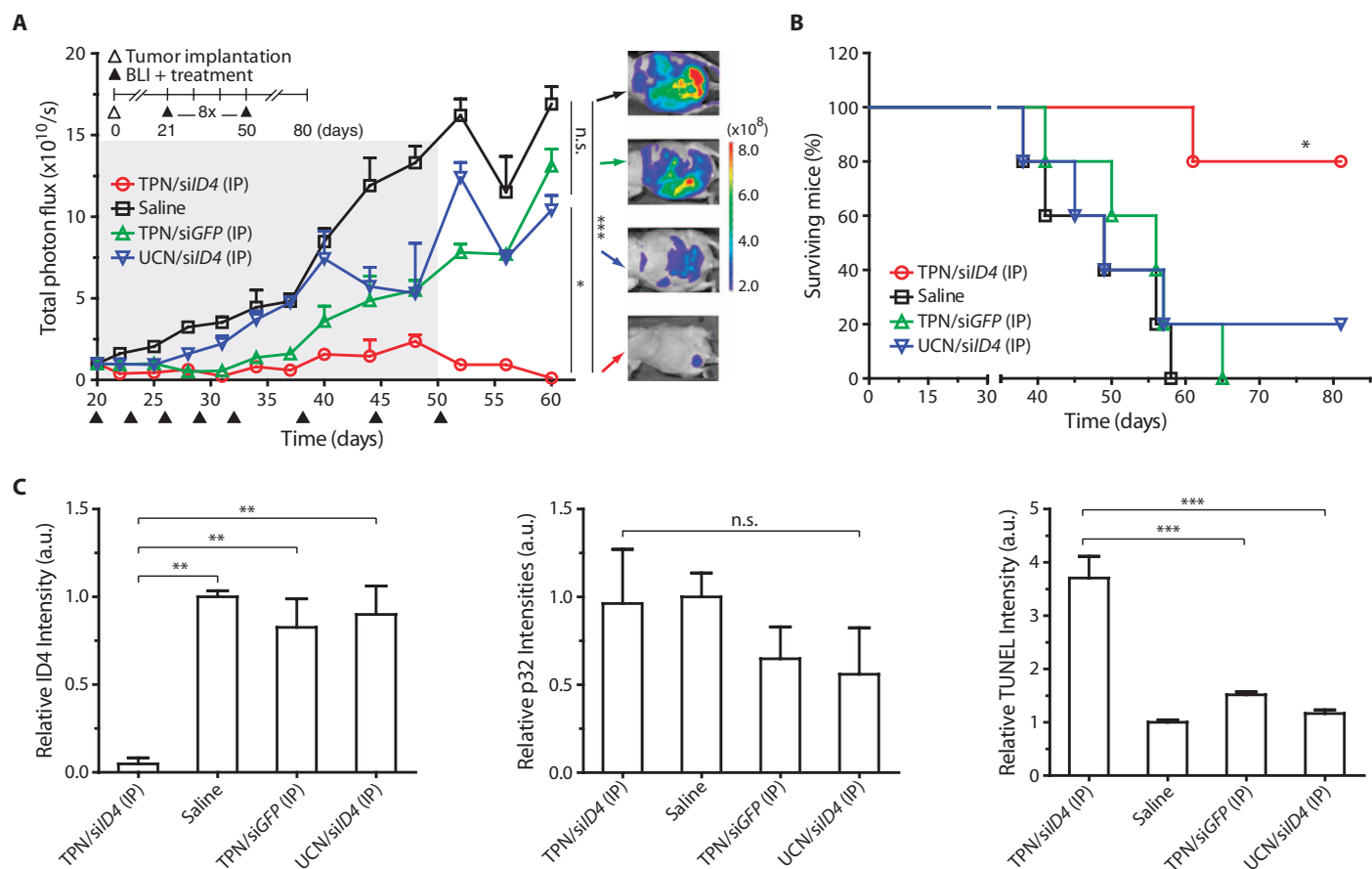


Fig. 7. Orthotopic tumor treatment with TPN/siD4. **(A)** Therapeutic efficacy of TPN in mice bearing disseminated orthotopic OVCAR-8 tumors. Mice were treated intraperitoneally every 3 days for 14 days and then once weekly thereafter for 3 weeks with TPN/siD4, saline, TPN/siGFP, and UCN/siD4 (arrowheads). Inset shows the experimental timeline. Total tumor burden was followed by bioluminescence imaging, with images from

day 60 shown. Data are means \pm SD ($n = 5$ mice per group). $*P < 0.05$, $***P < 0.001$, one-way ANOVA. **(B)** Kaplan-Meier plot of overall survival of the cohorts shown in (A). $*P < 0.05$ by log-rank (Mantel-Cox) test. **(C)** Quantification of ID4, p32, and TUNEL intensities from OVCAR-8 tumors on day 4. Intensity values are normalized to that of saline controls. Data are averages \pm SD ($n = 5$ per treatment group). $**P < 0.01$, $***P < 0.001$, one-way ANOVA.

scopic or histological signs of organ toxicity in the OVCAR-8 orthotopic tumor model (fig. S8C).

DISCUSSION

Current genome characterization efforts will eventually provide insight into the genetic alterations that occur in most cancers and may define new therapeutic targets. However, most epithelial cancers harbor hundreds of genetic alterations as a consequence of genomic instability. For example, whereas recurrent somatic alterations occur in a small number of genes in high-grade ovarian cancers, ovarian cancer genomes are characterized by multiple regions of copy number gain and loss involving at least 1825 genes. This genomic chaos complicates efforts to identify biologically relevant mutations critical for tumor maintenance.

To isolate which recurrent genetic alterations are involved in cancer initiation, tumor maintenance, and/or metastasis, functional assays can be performed after systematic manipulation of the candidate oncogenes. We have combined results from Project Achilles (2), a large-

scale screening effort to identify genes essential for proliferation and survival in human cancer cell lines with genome characterization of high-grade ovarian cancers. Using this approach, we identified an oncogene candidate, *ID4*, which was amplified in 32% of high-grade serous ovarian cancers. *ID4* is overexpressed in a large fraction of high-grade serous ovarian cancers, and ovarian cancer cell lines that overexpress *ID4* are highly dependent on *ID4* for survival and tumorigenicity. Expression of *ID4* at levels corresponding to those observed in patient-derived samples induced transformation of immortalized ovarian and FT epithelial cells. Our observations credential *ID4* as an ovarian cancer oncogene and suggest that *ID4* is one of a growing class of lineage-restricted transcriptional factor oncogenes in human epithelial cancers.

Like other ID family proteins, ID4 binds to and regulates the activity of E protein transcription factors such as transcription factors 3 and 12 (TCF3 and TCF12). Indeed, we found that ID4 regulates cell proliferation in part through its effects on p21^{WAF1/CIP1}. In addition, the transcriptional program induced by ID4 in ovarian epithelial cells was similar to that induced by HOXA9. Although it is clear that ID4 regulates the expression of many genes, we found that suppression of

HOXA9 abrogated *ID4*-induced cell transformation. *HOXA9* is known to regulate normal hematopoiesis by controlling differentiation, proliferation, and self-renewal (31), and overexpression of *HOXA9* or *HOXA9-NUP98* in hematopoietic precursors promotes the development of leukemias (32). In addition, *HOXA* genes have been shown to play essential roles in specifying regional differentiation of the Müllerian duct into oviduct, uterus, cervix, and vagina (33). Although previous work has shown that manipulation of *ID1*, *ID2*, or *ID3* expression induces compensatory changes in other *ID* family members, we found that expression of *ID4*, but not that of the other *ID* family members, induced cell transformation in ovarian epithelial cells. Because amplifications involving other *ID* family members have not been observed in ovarian cancers, it appears that *ID4* plays a unique role in the transformation of this tissue type. Collectively, these findings suggest that inappropriate overexpression of *ID4* may contribute to transformation by affecting both p21^{WAF1/CIP1} and *HOXA9* transcriptional programs that affect differentiation and proliferation programs in the ovary. Nevertheless, further mechanistic studies will be necessary to understand the biology of the new oncogene *ID4* and its downstream target, *HOXA9*.

To credential *ID4* as a therapeutic target in human ovarian cancer, we showed that a tandem peptide with both a membrane-translocating domain (TP) and a tumor-penetrating domain (LyP-1) condensed siRNA into nanocomplexes (TPN) that, upon systemic administration into ovarian tumor-bearing mice, penetrated the tumor parenchyma and silenced *ID4* in cells of interest (Fig. 5C). Treatment with TPN/si*ID4* suppressed the growth of ovarian tumors and significantly improved host survival. Previous approaches using inducible shRNAs to evaluate potential cancer targets were time-consuming and required extensive manipulation of cell lines in vitro (34, 35). Therefore, we favor this modular, tumor-specific, tumor-penetrating approach that delivers siRNA systemically. TPNs are biodegradable and did not elicit innate immune responses upon systemic delivery, enabling repeated administration for tumor regression studies.

Other methods for siRNA delivery include covalent coupling of siRNA to carriers, but this formulation process requires specialized chemically reactive siRNA and purification steps, obviating the modular “mix and dose” paradigm motivated by the volume of emerging genomic targets (36). Targeted siRNA delivery has been accomplished in specialized populations such as immune cells, neurons, and hepatocytes; however, these methods are not generalizable across tissue types. Elsewhere, siRNA has been targeted to tumors but not in a tumor-penetrating fashion (5). For example, tumor endothelium has been targeted using RGD-specific nanoparticles; however, this approach does not enable silencing in parenchymal tumor cells, such as those identified with genetic alterations in TCGA (1). Passive targeting (for example, the EPR effect) of siRNA in neutral liposomes, in complexes with CPP such as TAT, or transferrin-targeted cyclodextrin has also been described in mouse models and recently in humans (37, 38), but tumor penetration remained a challenge (39). Our TPN technology represents an important therapeutic paradigm for systemic targeting of undruggable proteins that are essential in parenchymal tumor maintenance.

Although we have focused on *ID4*, TPN delivery technology can be generalized to other cancers. Approximately 60 of 81 samples across 10 tumor types exhibit surface expression of p32 (20). Our modular siRNA delivery platform can leverage other homing peptides, such as iRGD, to expand the number and types of TPN-targeted parenchymal

tumor cells. More generally, ongoing cancer genome characterization efforts combined with comprehensive functional studies will narrow the number of potential cancer targets, but putative oncogenes will still require validation in vivo. TPN offers an efficient platform to verify which genes are critical to cancer initiation and maintenance.

Our studies demonstrate that TPN exhibits both specificity and pharmacokinetic properties that may provide a platform for human siRNA delivery. To reach this goal, further development is required, such as optimization of circulation time, assessment of efficacy in higher fidelity models of cancer, and reformulation to increase knock-down efficiency and reduce dosing. The path to translating TPN to the clinic will require multispecies toxicity testing, assessment of repeated dosing and potential immunostimulation, and evaluation of the prevalence of the tumor penetration pathway and p32 in human tumors and metastases.

In summary, we developed a targeted TPN capable of precisely delivering siRNA into the tumor parenchyma, and have combined this technology with large-scale methods to credential *ID4* as an oncogene target in ovarian cancer. Because large-scale efforts to characterize all cancer genomes accelerate, this capability illustrates a path to identify genes that are altered in tumors, validate those that are critical to cancer initiation and maintenance, and rapidly evaluate in vivo the subset of such genes amenable to RNAi therapies and clinical translation.

MATERIALS AND METHODS

Pooled shRNA library screen

shRNA constructs are available through Sigma-Aldrich. Genome-scale pooled shRNA screens in 102 cancer cell lines were performed with a lentivirally delivered pool of 54,020 shRNAs targeting 11,194 genes. Each cell line was infected in quadruplicate and propagated for at least 16 population doublings (2) (Supplementary Materials and Methods). The abundance of shRNAs relative to the initial DNA plasmids was measured by microarray hybridization and normalized with dChip and GenePattern modules before identifying essential genes.

Analysis of TCGA data

Copy number alterations were determined by TCGA on 489 primary ovarian tumors with Illumina 1M-Duo arrays, and recurrently amplified regions were analyzed by GISTIC 2.0 (40) with 99% confidence intervals (Supplementary Materials and Methods). For each amplified gene, the distribution of shRNA scores were converted to *z* scores, and shRNAs with a *P* < 0.05 were identified.

Cell culture

To determine the effects of shRNA on cell proliferation, we performed six replicate infections and determined viability 5 days later (Supplementary Materials and Methods). Transformation was determined by introducing *ID4* into IOSE-M (41) and FTSEC-M (FT secretory epithelial) cells (16) and performing anchorage-independent growth experiments or monitoring tumor formation by injecting cells subcutaneously into BALB/c immunodeficient mice (Charles River).

Gene expression profiling

Gene expression profiling was performed in triplicate on Affymetrix HG-U133A_2 GeneChips. Processed data from microarray experiments

were analyzed with GSEA (42) (Supplementary Materials and Methods).

Tumor-penetrating nanocomplexes

The tandem peptide library was synthesized by standard Fmoc (9-fluorenyl methoxycarbonyl) solid-phase peptide synthesis and purified by high-performance liquid chromatography (Supplementary Materials and Methods). Peptides were mixed with siRNA at a molar ratio of 20:1. Nanocomplexes were incubated over cell culture for 4 hours, and cells were harvested 48 hours later for characterization of gene knockdown. For competition with free LyP-1, unlabeled LyP-1 peptide was added for 1 hour before TPN treatment.

Therapeutic studies in mice

TPNs carrying *ID4*-specific siRNA were administered intravenously or intraperitoneally to NCr/nude mice bearing either subcutaneous OVCAR-4 or intraperitoneal disseminated OVCAR-8 tumors, respectively. Tumor burdens were monitored through caliper measurements or bioluminescence imaging (Supplementary Materials and Methods).

Statistical analysis

Unless otherwise indicated, one-way ANOVA was used (GraphPad). $P < 0.05$ was considered statistically significant. Fisher's exact test was used for tumor formation assays. Two-tailed Student's *t* test was used for pairwise comparisons. A log-rank test was performed for animal survival studies.

SUPPLEMENTARY MATERIALS

www.sciencetranslationalmedicine.org/cgi/content/full/4/147/147ra112/DC1
Materials and Methods

Fig. S1. Amplification and overexpression of *ID4* in ovarian cancers.

Fig. S2. *ID4* regulates *HOXA9* gene activity.

Fig. S3. *ID4* regulates expression of *HOXA9* in ovarian cancer cells.

Fig. S4. In vitro characterization of TPNs.

Fig. S5. TPN-mediated suppression of *ID4* in p32-expressing ovarian cancer cells.

Fig. S6. Biodistribution of TPN.

Fig. S7. Therapeutic effects of *ID4* suppression in ovarian tumor-bearing mice.

Fig. S8. Lack of toxicity of *siID4*.

Table S1. List of amplified genes with at least one shRNA that score with a $P < 0.05$.

Table S2. Sequence and characterization of tandem peptides.

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