

Chemical engineering of therapeutic siRNAs for allele-specific gene silencing *in vivo* in CNS

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Abstract

Small interfering RNAs (siRNAs) are a new class of drugs, exhibiting sequence-driven, potent, and sustained silencing of gene expression *in vivo*. We recently demonstrated that siRNA chemical architectures can be optimized to provide efficient delivery to the CNS. Many genetically-defined neurodegenerative disorders are autosomal dominant favoring selective silencing of the mutant allele. In some cases, successful targeting of the mutant allele requires targeting of a single nucleotide polymorphism (SNP) heterozygosity. Using Huntington's disease as a model, we demonstrate allele-specific RNAi-based silencing of gene expression *in vivo* and in neurons differentiated from HD patient-derived iPSCs. A series of *in vitro* screens, with chemical and thermodynamic optimization, identified compounds with >50-fold selectivity for the mutant HD-causing allele, based on a single nucleotide difference. The optimized compound exhibits selective silencing of mutant huntingtin (HTT) protein in patient derived cells and throughout the HD mouse brain, providing a demonstration of SNP-based allele-specific RNAi silencing of gene expression *in vivo* in the CNS. The ability to target a disease-causing allele

using RNAi-based therapies could be applied to a wide range of dominant CNS disorders, where maintenance of wild-type expression is essential.

Introduction

Oligonucleotide therapeutics is a novel class of drugs advancing the treatment of a wide range of diseases^{1,2}. The clinical success of these compounds is predicated on an optimized chemical scaffold supporting delivery to the tissue of interest; the sequence can then be modified to target other genes of interest.^{3,4} One class of oligonucleotide therapeutics is small interfering RNAs (siRNAs). Upon loading into the RNA-induced silencing complex (RISC), siRNAs cleave complementary mRNAs leading to degradation of the disease-causing mRNA and preventing protein expression⁵⁻⁸.

Stabilization of the siRNA is critical for sustained efficacy. A single injection of a multi-valent, GalNac-conjugated, fully-modified siRNA safely supports target silencing in the liver for up to 6-12 months^{1,9-11}. Endosomal entrapment of oligonucleotides creates an intracellular depot and provides a continuous supply of siRNA for RISC loading¹², explaining the long duration of effect. A combination of sugar, backbone, and terminal phosphate modifications¹³⁻¹⁸ stabilizes the compound in the endosomal compartment and is required to ensure durable gene silencing *in vivo*.

Recently, we developed an optimized siRNA chemical architecture that achieves broad distribution and long-lasting silencing in the brains of rodents and non-human primates. The divalent, chemically modified (Di-siRNA) exhibited delayed cerebrospinal fluid (CSF) clearance and efficient neuronal internalization¹⁹. Furthermore, a single intracerebroventricular (ICV) injection of a Di-siRNA targeting the *HTT* mRNA produced widespread silencing of total *HTT* for up to six months.

Huntington's disease (HD) is an autosomal dominant neurodegenerative disorder caused by the expansion of the CAG repeat region of the Huntington (*HTT*) gene. Reducing huntingtin protein (*htt*) expression is a therapeutic approach^{20,21} that has shown promise in multiple disease models²²⁻²⁵. Recent evidence indicates that wild-type huntingtin is might be important for normal neuronal function^{26,27}, therefore, selective silencing of the mutant isoform might be advantageous. Moreover, a large phase III clinical trial evaluating the efficacy of a non-selective antisense oligonucleotide (ASO) targeting HTT was recently halted prematurely due to negative effects^{28,29}. ASO-HTT treated patients showed an increase in ventricular volume, increased expression of neurofilament light chain (NfL), and a decrease in functional performance readouts. While the negative outcome is likely related to non-specific inflammatory effects of ASOs, the potential impacts of non-selective reduction of HTT cannot be ruled out. Like HD, many genetic disorders are dominant³⁰, and silencing of the mutant, but not the wild type allele, may be preferable³¹⁻³⁴. One strategy for allelic discrimination targets SNP heterozygosities^{35,36} present in normal and disease-causing genes^{35,36}. While theoretically ideal, SNP-based allele-specific silencing using chemically modified siRNAs has never been demonstrated in brain.

Multiple reports describe the *in vitro* development and validation of unmodified siRNAs that effectively discriminate between mutant and wild-type alleles based on a single nucleotide³⁵⁻⁴². However, the degree of discrimination achieved was 10-fold or less⁴³, which may not translate into effective discrimination *in vivo* where siRNA accumulation varies regionally and temporally⁴⁴. In addition, most *in vitro* studies were performed using non-modified or partially modified siRNAs, which are unstable and do not distribute widely *in vivo*. Full chemical stabilization is essential for *in vivo* efficacy⁴⁵ but can significantly affect guide strand

thermodynamics⁴⁶. Thus, the chemical modification of siRNAs changes their target recognition and discrimination properties, altering their SNP targeting profile.

Using a fully chemically modified siRNA scaffold, we performed a systematic series of screens targeting two SNPs, rs362307 and rs362273, that are frequently heterozygous in HD patients^{35,36}. For rs362273, we identified one siRNA exhibiting greater than 50-fold discrimination and confirmed the selectivity of this siRNA in human neurons differentiated from HD patient iPSCs. In the CNS-active (Di-siRNA) conformation, the optimized siRNA showed widespread distribution in a transgenic mouse model of HD (BACHD)⁴⁷ and maintained allele-specific silencing in all brain regions, independent of dose or level of siRNA accumulation. These results demonstrate RNAi-based allele-specific silencing using fully modified siRNA *in vivo*. The approach—demonstrated for HD—can be applied to numerous other genetic disorders where allele-specific mutant gene silencing is necessary.

Results

Multiple SNPs in the *htt* gene are frequently heterozygous in patients^{35,36,48}. Here, we focus on two SNPs in the *htt* gene (Supplementary Figure 1): rs362273 in exon 57, which is heterozygous in 35% of HD patients, and rs362307 in exon 67, which is heterozygous in 48% of HD patients^{35,45}. One of the challenges for *in vivo* evaluation of SNP-selective siRNAs is access to *in vivo* models. For *in vivo* evaluation, we chose SNP rs362273, which is present in BACHD mice. The surrounding region is conserved in the mouse *htt* gene creating a model where the mutant (human, expanded CAG repeat) and the wild-type (mouse) HTT differ by a single nucleotide at the SNP site. Because the mutant allele has a long CAG repeat, the mutant and wild-type HTT proteins can be separated by western blot and evaluated simultaneously for selective silencing.

Primary screen generates siRNAs with moderate SNP-based discrimination

We designed a panel of twelve chemically stabilized siRNAs, overlapping SNP rs362273 of the huntingtin mRNA (Figure 1). Supplementary Table 1 shows the sequences and chemical modification patterns of all compounds used in the study. Alternating 2'-O-methyl (2'-OMe) and 2'-deoxy-2'-fluoro (2'-F) replaced the riboses, and terminal backbones were modified with phosphorothioates (PS); the sense strand was conjugated to cholesterol (Figure 1a). We refer to these siRNAs as fully chemically modified. Cholesterol conjugated, chemically stabilized siRNAs are efficiently internalized in all cell types through endocytosis and lysosomal entrapment^{49,50}. The cholesterol-driven self-delivery mechanism is analogous to the mechanism of *in vivo* uptake and requires extensive stabilization. Therefore, the same chemical scaffold, with the same pattern of 2'-F, 2'-OMe, and PS modifications, but without cholesterol, can be re-synthesized in the divalent, CNS-active configuration for a seamless transition between *in vitro* and *in vivo* experimental systems¹⁹.

For evaluation of on-target and non-target efficacy, we used two luciferase-based reporter plasmids containing a 40-nucleotide portion of the huntingtin mRNA surrounding SNP site rs362273⁴⁵. Cells were transfected with either the 2273-A or 2273-G reporter plasmid. The next day, the cholesterol-conjugated siRNA was added. Figure 1b shows the results of the primary screen. The silencing of target and non-target reporters varied significantly depending on siRNA sequence and position of the SNP heterozygosity. For example, the siRNA with the SNP in position 3 showed efficacy but little discrimination, while those with the SNP in positions 12 and 13 showed limited efficacy.

For compounds with the highest on-target efficacy and discrimination, we evaluated the efficiency of target and non-target silencing at multiple concentrations and calculated IC50 values

(Figure 1c). Two siRNAs, SNP4 (SNP in position 4) and SNP6 (SNP in position 6), exhibited potent target gene silencing (IC₅₀ ~38nM and 24nM correspondingly) and demonstrated ~20-fold (SNP4) and ~7-fold (SNP6) discrimination against the non-target reporter based on a single mismatch. We proceeded to optimize these two candidates.

The SNP targeting siRNAs is two to four-fold less potent than the most active pan-targeting siRNAs in the native genomic context

Extensive screening across the whole gene can identify highly potent pan-targeting siRNAs but the sequence space available for identifying SNP targeting compounds is limited. By screening hundreds of siRNAs, we have previously identified a highly active non-selective siRNA^{HTT} targeting HTT⁵⁰. HeLa cells express endogenous HTT and have the A isoform at the rs362273 SNP site. Therefore, we can compare the potencies of SNP- and non-selective siRNAs against the endogenous target. We evaluated the efficacy of SNP4, SNP6, and siRNA^{HTT} in HeLa cells at seven concentrations (Supplementary Figure 2). In passive uptake, the corresponding IC₅₀ values for SNP4, SNP6, and siRNA^{HTT} were 302nM, 149nM, 73nM, indicating that limiting the targeted sequence space by targeting a SNP resulted in at least a two-fold reduction in potency.

Introduction of a secondary mismatch enhances SNP-based discrimination

The level of siRNA accumulation *in vivo* varies significantly between different brain regions and over time. Therefore, a single mismatch at the optimal position (SNP6) likely provides insufficient discrimination to support allele-specific silencing *in vivo*. In the context of a fully modified guide strand, a single mismatch introduces only a small thermodynamic disturbance, and indeed in certain positions, a single mismatch between the siRNA and the target mRNA can enhance efficacy⁵¹. We measured the impact of the mismatch in position six on the Tm of the guide strand/RNA substrate (Supplementary Figure 3). In the context of the full-length guide, the impact

of the mismatch on stability was 4°C (85.4°C to 81.4°C), which is not biologically significant. In the context of a 13-mer substrate (comprising the RISC core interactions), the impact of the mismatch was greater (5.1°C—68.5°C to 73.6°C). However, in both cases, the affinity of the modified guide strand for the target is high. The likely explanation for the observed discrimination is a disruption of the local architecture of the siRNA-target duplex, which interferes with the ability of RISC to form the active conformation and disrupts scanning and recognition of the target by the seed region (positions 2-8)⁵²⁻⁵⁴.

Introducing a secondary mismatch⁴⁵ can enhance SNP-based discrimination by modulating guide strand affinity for the target. In the context of a SNP heterozygosity, siRNAs with a secondary mismatch (Figure 2a) have a single mismatch to the target isoform and two mismatches to the non-target isoform. The second mismatch can significantly reduce the affinity of the guide strand for the double-mismatched non-targeted isoform, resulting in a loss of silencing⁴⁵. Using the two best siRNAs from the primary screen, SNP6 (Figure 2b) and SNP4 (Supplementary Figure 4), we performed a secondary mismatch screen. Introducing an intentional mismatch to the SNP6 siRNA at position 11 enhances the level of target/non-target discrimination to more than 50-fold (Figure 2b).

Ago2 uses the seed region for primary target recognition⁵². Additional complementarity throughout the rest of the guide stand can compensate for an imperfect seed. A secondary mismatch outside of the seed region minimally impacts target silencing but reduces the silencing of the non-targeted mRNA isoform (Figure 2a). To quantify the differences in target and non-target recognition, we evaluated compounds with secondary mismatches in positions 11, 14, or 16 in a 7-point dose-response (Figure 2b). None of these secondary mismatches affected on-target silencing (IC50s ~20-30 nM). The impact on non-target silencing varied, increasing discrimination

to greater than ~50-fold (Figure 2b). The position of the best secondary mismatch is sequence-dependent; a secondary mismatch in position 7 improves discrimination when the primary mismatch was at position 4 (Supplementary Figure 4b), and a position 11 secondary mismatch was better when the primary mismatch was at position 6 (Figure 2b). Therefore, the exact combination of primary and secondary mismatches needs to be optimized individually for each sequence. For further optimization, we selected compound SNP6-11, which carries two mismatches and showed good on-target activity (IC₅₀ ~28 nM), with more than 58x discrimination against the non-targeted isoform (IC₅₀ >1500 nM).

Chemical modifications at the mismatch positions affect potency and discrimination

While both 2'-F and 2'-OMe stabilize the siRNA, the impact of 2'-OMe modification on stability is significantly more pronounced⁵⁵. Thus, increases in 2'-OMe content improve siRNA stability and enhance duration of effect *in vivo*^{16,56}, but extensive 2'-OMe modifications can position and sequence-specifically decrease activity⁵⁷. Therefore, most siRNAs designed for *in vivo* and clinical applications use sequence-optimized patterns of 2'-OMe and 2'-F modifications⁴. Successful development of SNP targeting siRNA requires an understanding of how the chemical modification pattern around the mismatch affects efficacy and discrimination. 2'-OMe and 2'-F affect guide strand/target interactions differently. 2'-F strengthens target interaction substantially more than does 2'-OMe^{58,59}. An increase in 2'-F content, particularly around the mismatch site, is expected to increase local affinity of the siRNA for its target, reducing the impact of a mismatch. Meanwhile, the bulkier 2'-OMe might increase discrimination by reducing the silencing of the non-targeted isoform.

A panel of compounds with different combinations of 2'-OMe and 2'-F modifications near the primary (position 6) and secondary (position 11) mismatch positions were synthesized and

tested. Changes in the chemical modification pattern around the mismatch sites affected potency and discrimination (Figure 3a). Figure 3b-e shows the dose responses for the most informative configurations. Placing 2'-F modifications flanking the primary site (SNP fm6-11, Figure 3c) maintains on-target potency (IC₅₀ ~17nM vs 28nM—see Figure 4b) but eliminates discrimination. Increased 2'-OMe content around the secondary mismatch (SNP 6-m11) (Figure 3d) reduces on-target silencing (IC₅₀ ~58nM). Combining 2'-F modifications around the primary SNP site with a methyl rich secondary mismatch region (SNP f6-m11, Figure 3e) restores on-target activity (IC₅₀ ~22nM) and maintains discrimination. These data suggest that both structural and chemical contexts near the mismatch affect siRNA activity and selectivity. Therefore, SNP selectivity must be optimized in the same chemical context intended for *in vivo* use.

The optimal SNP-discrimination scaffold is sequence-specific

Having identified a highly selective siRNA targeting HD SNP rs362273, we tested whether this approach could be applied to other sequences. We repeated the same workflow, this time targeting HD SNP rs362307, which is heterozygous in 48% of HD patients.⁴⁵ This SNP is not present in current HD mouse models⁶⁰, precluding *in vivo* evaluation.

The primary screen (Supplementary Figure 5a) identified only one siRNA with acceptable activity (IC₅₀ target ~35nM) and discrimination (IC₅₀ for non-target ~438nM) (SNP3, Supplementary Figure 5b). A secondary mismatch screen demonstrated that both the position and the identity of the mismatch affected potency and selectivity (Supplementary Figure 5c). Introduction of a secondary mismatch at position 8 increased selectivity (Supplementary Figure 5d) whereas a secondary mismatch at position 5 increased selectivity, but also decreased on target potency (Supplementary Figure 5e). In addition to the position, the type of mismatch was important. For example, the SNP3-7C (Supplementary Figure 5f) compound had little activity

toward the non-targeted allele ($IC_{50} > 1500\text{nM}$) but lost up to 5-fold in target silencing efficacy ($IC_{50} \sim 400\text{nM}$) relative to the single mismatch SNP3 compound. SNP3-7G (Supplementary Figure 5g) shares the same mismatch positions as SNP3-7C but differs in sequence at the secondary mismatch position. SNP3-7G showed the same efficacy against the targeted allele as SNP3-7C but did not discriminate between alleles. Consistent with the idea that the wobble base-pair is less structurally disruptive⁴³, the C-U mismatch discriminates better than the G-U mismatch⁶¹ (SNP3-7C vs. SNP3-7G, Supplementary Figure 5f,g). The optimal scaffold SNP3-5G (Supplementary Figure 5e) exhibited effective target silencing ($IC_{50} \sim 92\text{nM}$) and efficient discrimination (non-targeted $IC_{50} > 1500\text{nM}$). These data suggest that both the chemical and sequence context of the mismatches influence efficacy and discrimination, requiring individual optimization.

Allele-selective lowering of mutant HTT protein in human neural stem cells and in stem cells differentiated into neurons

To determine if SNP6-11 could discriminate between wild-type and mutant huntingtin in a native human context, we treated human neural stem cells (NSCs) with SNP6-11 siRNA. The NSCs were created from induced pluripotent stem cells (iPSCs) with 109 CAG repeats. We performed DNA sequencing and determined that these cells were heterozygous at rs362273. There are three major huntingtin haplogroups, A, B and C⁶². A majority of HD chromosomes belong to the A haplogroup, representing targetable SNPs associated with HD. In contrast, the C haplogroup is overrepresented on normal chromosomes⁶². Evidence from other SNP heterozygosities suggests that the parental CAG109 cell line belonged to the A or B haplogroups⁶³⁻⁶⁵. Both A and B haplogroups have an A on the mutant isoform at the rs362273 SNP site; since we had established

heterozygosity, we reasoned that the non-HD chromosome would belong to haplotype C and have a G at the SNP site.

After treatment with siRNA, we measured the levels of wild-type and mutant HTT protein by SDS-PAGE and western blot. The non-selective siRNA^{HTT} lowered both wild-type and mutant HTT protein (Figure 4a). In contrast, when compared to a non-targeting control (NTC) at 3.0 μ M, SNP6-11 produced a dose-responsive lowering of mutant HTT at 1.5, 2.0, and 3.0 μ M (Figure 4a) with no corresponding reduction of wild-type HTT (Figure 4a). This data indicates that based on a single nucleotide difference and in the presence of two native human *htt* alleles, the SNP 6-11 siRNA selectively lowers mutant HTT (A) without affecting the wildtype protein (G).

Next, we asked if SNP6-11 could reduce mutant HTT protein selectively in a disease-relevant cell type. HD109 iPSCs were differentiated to cortical neurons. Western blot analysis showed that the neuronal marker β III-tubulin was present in cell lysates and immunofluorescent labeling of parallel wells revealed a high percentage of β III-tubulin positive cells, confirming neuronal enrichment (Supplementary Figure 6a,b). siRNAs were added for 5 days continuously, starting on differentiation day 43, as described in Methods. Compared to NTC at 3.0 μ M, SNP6-11 at 2.0 and 3.0 μ M significantly lowered mutant HTT protein by 35 and 38%, respectively (Figure 5a). In contrast, levels of wild-type HTT protein were unchanged (Figure 5b). At 3.0 μ M, non-selective siRNA^{HTT} lowered both mutant and wild-type HTT protein by 80% and 78% (Figure 5a,b). Like in HeLa cells (Supplementary Figure 2), the SNP 6-11 siRNA shows efficient discrimination between the two alleles but lower efficacy when compared to the non-selective siRNA^{HTT}.

SNP targeting siRNA compounds selectively silence mutant huntingtin in the mouse brain

The next step was to evaluate whether the SNP6-11 compound, targeting rs362273, can discriminate between mutant and wild-type huntingtin *in vivo*. The BAC-HD mouse model expresses wild-type mouse huntingtin and overexpresses a human HD transgene⁴⁷. The sequence overlapping rs362273 is conserved between human and mouse genes, with the mouse *htt* having an A, and the transgene a G at the SNP position (Figure 6a). Therefore, we can evaluate selective SNP-based silencing in this model. For *in vivo* delivery, the SNP6-11 compound was synthesized in a divalent siRNA configuration (Figure 6b), which supports robust silencing in the mouse and NHP brain¹⁹. Supplementary Table 1 shows sequence and chemical modification patterns used.

We administered 225 µg (10 nmol, 5 nmol/ventricle) of divalent SNP6-11 into the lateral ventricles of BACHD mice. PBS and a chemically matched divalent NTC siRNA were used as negative controls. One-month post-injection, the levels of mutant human and wild-type mouse huntingtin protein were measured in the cortex, striatum, hippocampus, and thalamus. There were no significant differences between PBS and NTC, indicating that the divalent siRNA chemical scaffold alone does not change expression of either the wild type or the mutant huntingtin alleles (Figure 6c,d). The animals treated with SNP6-11 showed >85% silencing ($P < 0.0001$) of the mutant protein with no change in levels of the wild-type HTT (Figure 6c,d). Divalent siRNA accumulation in different brain regions varies⁶⁶, but selective silencing of mutant huntingtin protein is sustained throughout the brain, indicating that more than 50x discrimination *in vitro* is sufficient to support selective silencing *in vivo*. Selective mutant *htt* silencing did not change the levels of the neuron-specific protein Dopamine- and cAMP-regulated phosphoprotein (DARPP32; Supplementary Figure 7a). Glial fibrillary acid protein (GFAP), a marker of astrocytes, was also unchanged, except in the striatum (Supplementary Figure 7b). Levels of ionized calcium-binding adaptor molecule (Iba1), which localizes to microglia, were not different between treatment groups

(Supplementary Figure 8), indicating that exposure to the divalent siRNA did not elicit significant microglial activation at the times tested.

To ensure that selective allele-specific silencing *in vivo* also occurs at a high siRNA dose, we injected the second cohort of animals with 450 μ g (20nmol, 10nmol/ventricle) of divalent siRNA, the maximum deliverable dose for this chemical class. Like the 225 μ g injection, selective silencing of mutant *HTT* occurred throughout the brain (Supplementary Figure 9). At one month, the time of greatest oligonucleotide accumulation, mutant HTT approached the lower detection limit (> 99% silencing, P<0.0001) with no decrease in wild-type huntingtin, indicating that SNP6-11 can support selective silencing at a wide range of doses throughout the brain over time. We expect this higher dose to maintain silencing for longer than 6 months⁶⁶.

Here we demonstrate that using a fully chemically modified siRNA, we can achieve selective silencing of mutant *HTT* *in vivo*, based on a SNP heterozygosity. We expect this approach to be generalizable to other autosomal dominant disorders where expression of the wild-type allele is essential.

Discussion

Allele-specific modulation of gene expression in the CNS promises to preserve normal function and enable disease-modifying treatment for many neurodegenerative disorders. Here, we demonstrate that fully chemically modified, therapeutically translatable siRNAs targeting SNP heterozygosities support allele-specific discrimination in human neurons derived from human patient iPSCs and *in vivo* in mice. Through repeated targeted screens and chemical optimization, we identified SNP-selective siRNAs that achieved more than 50x discriminative power in a cell-based assay and confirmed selective silencing of the mutant *HTT* allele (>85%) throughout the

brain in a mouse model of HD. Sequence context and chemical composition affected selectivity; optimization for each SNP heterozygosity should be performed in the chemical configuration intended for *in vivo* use. Our findings provide a roadmap for identifying allele-specific therapeutic siRNAs for other disorders in which expression of the normal allele is essential and indicate that compounds with 50x discrimination in cell culture support robust allele-selective silencing *in vivo*.

For SNP-discriminating siRNAs, the sequence space available for screening is limited to approximately 16 bases immediately surrounding the SNP position. The placement of mismatches proved crucial for siRNA selectivity. For the rs362273 A isoform, mismatches in positions 6 and 11 showed the best potency to discrimination ratio. Whereas for rs362307 U isoform, positions 3 and 7 were optimal. These findings suggest that for maximum discrimination, the positioning of the SNP heterozygosity and of the second mismatch is dependent on the target sequence. Thus, the position, identity, and modification pattern of both primary and secondary mismatch need to be optimized individually for different SNP sites.

Although we identified SNP-selective siRNAs against both SNP sites evaluated, similar success might prove challenging for some SNP heterozygosities, particularly if the sequence around the SNP site is GC rich⁶⁷. siRNAs vary vastly in their ability to silence their targeted mRNAs, with EC50 values ranging from the low pM to the low nM range. The efficacy of an siRNA is affected by the ability to enter RISC and the efficiency with which the loaded RISC complex can find the target, adopt the right conformation for cleavage, and release the resulting product^{54,68}. As the sequence space available for screening of SNP-selective compounds is limited, their potency could be lower than the best non-selective compounds for the same target. For example, the potency of both SNP-targeting compounds was less than siRNA^{HTT}, the previously identified non-selective HTT-targeting siRNA^{19,50}. Consequently, therapeutic translation of SNP-

selective compounds may require higher doses or more frequent dosing to achieve silencing comparable to their non-selective counterparts.

Chemical context near the SNP site affects selectivity. The introduction of 2'-F modifications increases the local oligonucleotide affinity⁵⁸. Placing this modification near the SNP position increases the silencing of both alleles but reduces discrimination (Figure 4). In contrast, 2'-OMe modifications reduce siRNA affinity to the target and, in many positions, are not sterically well tolerated. Manipulating affinity throughout the siRNA affected target discrimination. Introduction of bulky 2'-OMe modifications at mismatch sites increased discrimination while higher affinity 2'-F modifications decreased it. By optimizing sequence, structure and chemical modification pattern, we identified multiple functional compounds with SNP-discriminating properties; however minor changes in the chemical modifications (e.g. a change from 2'-OMe to 2'-F in a single position) impacted both on-target activity and discrimination (Figure 3). Therefore, optimization for efficacy and selectivity must be performed in the context of the modification pattern intended for *in vivo* use. Further exploration of an expanded range of chemical modifications by including, for example by bulkier entities like MOE (2'-methoxyethyl), or highly flexible UNAs (unlocked nucleic acid) might eliminate the need for structural mismatches.

The level of discrimination achieved in the cell-based assay with the rs362273 targeting SNP6-11A siRNA, (~58x difference in IC50 values) was sufficient for allele selective silencing *in vivo* in CNS. Treatment in mice achieved efficient silencing of mutant HTT protein in all brain regions without impact on wild-type HTT protein, despite varied siRNA accumulation throughout the brain. The thalamus accumulated ~ 10-15x more siRNA than did the cortex (4,24). All brain regions demonstrated effective allelic huntingtin lowering. Nonetheless, the pharmacokinetic dose range of selective silencing in brain did not apply to non-brain tissues. The systemic administration

of GalNac modified siRNA supports about a ten-fold higher accumulation in the liver compared to brain. The increased accumulation in liver dampened the selectivity of siRNA6-11. To recover oligonucleotide selectivity, introduction of an additional stereo-constrained vinyl phosphonate backbone next to the mismatch site was necessary to further enhance discrimination and maintain selectivity *in vivo* in the liver⁶⁹. If the same accumulation in the brain as in the liver can be achieved, additional chemical enhancement of the siRNA might be necessary.

In HD and other expanded repeat-associated diseases, the causative mutation is not amenable to direct targeting. Therefore, a panel of SNP-targeting compounds must be developed. Studies of the HD patient population and targetable SNPs^{48,64,70,71} suggest that 80-85% of HD patients can be treated with panels of 2-4 SNP heterozygosities^{35,36,64,71}. Up to 40% of patients of European descent are heterozygous at rs362273³⁶ and the A isoform is more common on HD chromosomes⁶⁵, we have successfully targeted the A isoform, but to treat 40% of patients both isoforms would need to be targeted. If the allele selective silencing of HTT becomes essential for clinical disease modification then, the linkage between the CAG repeats and the SNP site heterozygosity would need to be established individually for eligible patients using long read sequencing⁷².

Materials and Methods

Cell treatment: reporter assay

HeLa cells grown and maintained in DMEM (Gibco ref.# 11965-092) with 1% penicillin /streptomycin and 10% heat inactivated fetal bovine serum (FBS). Three days prior to treatment, two 10cm² dishes are plated with 2x10⁶ HeLa cells and grown overnight. The following day,

DMEM is replaced with OptiMEM (Gibco ref.# 31985-070) and 6ug of reporter plasmid⁴⁵ is added to cells using lipofectamine 3000 (Invitrogen ref.# L3000-015), following the manufacturer's protocol. Cells are left in the OptiMEM/lipofectamine overnight to allow for maximum reporter plasmid transfection. The following day, cells are ready for treatment.

For screening purposes, siRNAs are prepared in OptiMEM at 3 μ M concentration. 50 μ L of 3 μ M siRNA is then added to 96-well white wall clear bottom tissue culture plate, in triplicate, for each reporter plasmid. For a dose response curve, 800 μ L of 3 μ M siRNA is prepared in OptiMEM, and a 1:1 dilution series is performed six times for seven concentrations of each siRNA: 3 μ M, 1.5 μ M, 0.75 μ M, 0.375 μ M, 0.1875 μ M, 0.09375 μ M, 0.046875 μ M. 50 μ L of each siRNA dilution are then added to each white wall clear bottom 96 well plate in triplicate. After the desired amounts of siRNA were added to the plate, 50 μ L per well of HeLa cells transfected with reporter plasmids were added, resuspended in DMEM with 6% heat inactivated FBS at 0.15 \times 10⁶ cells/mL, bringing maximum final siRNA concentration to 1.5 μ M.

After 72 hours of treatment (100% confluency) cells were washed twice with PBS, then lysed with 20 μ L 1x passive lysis buffer from Promega dual-luciferase assay system kit (Promega ref.# E1960) and placed on the shaker for 15-20 minutes. Luminescence was read after adding 50 μ L Luciferase Assay Reagent II (Promega kit), then read a second time after addition of 50 μ L/well of stop and glow reagent. Luminescence values were normalized to untreated controls and graphed on a log scale.

Stem Cells

Experiments were performed with oversight of Human Embryonic Stem Cell Research Oversight Committee (ESCR Committee) through the Partners (now MGB) Institutional Biosafety Committee (PIBC) at Partners Healthcare (ESCR#: 2015-01-02 and PIBC Reg#

2017B000023). Human induced pluripotent stem cells (iPSCs) CS09iCTR-109n4 (CS vial ID: 1034860) and CS09iCTR-109n5 (CS vial ID: 1034589) hereafter referred as HD109 (clone n4 and n5) described by (Mattis VB et al., 2015)⁷³, were acquired from Cedars-Sinai Regenerative Medicine Institute Induced Pluripotent Stem Cell Core. Cells were grown in mTeSR™ Plus kit (StemCell Technologies, 100-0276) on plastic plates coated with Matrigel (Corning, 354277) Cell lines were karyotyped after in-house expansion by Cell Line Genetics using array comparative genomic hybridization (aCGH, Agilent 60K Standard). Limited chromosomal imbalances in non-critical genes were reported: a single deletion of 460kb in chromosome 2, cytoband q22.1 (*LRP1B*) in HD109 n4, while HD109 n5 displayed an additional 90kb deletion in chromosome 14, cytoband 24.3 (*SPTLC2*). The DNA sequence of HTT at SNP RS362273 (hereafter named SNP2273) was determined via Sanger sequencing by Genewiz, Inc. using primers previously published⁴⁵ and shown to be heterozygous (G/A) for HD109.

Neural Stem Cells (NSCs)

NSCs were differentiated as described⁷⁴. NSCs were maintained in NSC Medium without 1% KnockOut serum replacement and plated on 48-well (Corning, 3548) plastic plates coated with 0.1% Gelatin for siRNA experiments.

siRNA treatment of NSCs and Neurons

For NSCs, normal growth medium was replaced with NSC medium without antibiotics and containing cholesterol-tagged siRNAs targeting total HTT (HTT10150), allele specific HTT siRNAs, or a non-targeting control (NTC) diluted to indicated 1x in 48 well plates. NSCs were treated for 5 days with a complete medium change at day 3 with medium containing fresh siRNAs at 1x concentration. For human neuron cultures in 48 well plates, on differentiation day 42 NMM was switched 250 µl/well to NMM without antibiotics/mitotic inhibitors 24 hours prior

to treatment. On Day 0 of treatment (differentiation day 43), 200 μ L of NMM was removed from each well leaving 50 μ l remaining, and 250 μ L/well of complete medium containing siRNAs at 2x concentrations were introduced bringing the volume to 300 μ l (0.83 dilution factor; final concentration of siRNAs, 1.66x). After 24 hours of treatment, additional 200 μ L of NMM without antibiotics/mitotic inhibitors was added to each well brining the volume to 500 μ l (0.6x dilution factor; final concentration of siRNAs, 1x) and incubated for 4 more days (total 5 day treatment). Concentrations indicated in Figure 4 those of 4 day incubation (1x). 30 μ l of lysis buffer (50 mM Tris, pH 7.4, 250 mM NaCl, 1% Nonidet P-40, 5 mM EDTA) containing protease inhibitors per well was used to lyse cells for western blot analysis.

Stereotactic brain injections into the lateral ventricle

BAC97 HD⁴⁷ mice were bred and genotyped via PCR to validate presence of the BAC97 transgene. At approximately 8 weeks of age, BAC97 females were anesthetized with tribromoethanol at 284mg/kg weight. Mice were subsequently placed in a stereotaxis and hair was removed from scalp using glue. The injection site was cleaned and prepared for surgery with three applications of betadine and ethanol swabs. An incision was made at the top of the scalp and the bregma was located as a reference point for injection sites. The needle was placed +/- 0.8mm mediolateral and 0.2mm posterior to the bregma on both sides of skull for bilateral injection. Holes were drilled and the needle was lowered 2.5mm for placement into the lateral ventricle. 5uL of 4000 μ M siRNA suspended in PBS was injected into the right lateral ventricle at a rate of infusion of 500nL/minute, followed by an identical injection into the left ventricle, for a total injection of 10 μ L/40nmols siRNA. Needles were left in place for one minute following the end of infusion to avoid backflow. Upon removal from stereotaxis, the incision was closed

with staples and Meloxicam SR was administered for analgesia. Mice recovered on a heating pad until awake and sternal.

Tissues were harvested one-month post-injection. Mice were euthanized with isofluorane and brains were removed and cut into 300 micron sections using a vibratome submerged in PBS at ~ 0°C. Sections were suspended in PBS on ice, where 2mm punches were taken using a sterile disposable biopsy punch (Integra Militex REF#33-31-P/25) from 3 sections bilaterally for each brain region: striatum, medial cortex, posterior cortex, hippocampus, and thalamus. Punches from one side of the brain were flash frozen in liquid nitrogen for subsequent protein analysis. Punches from the opposite side of the brain were saved in RNA later for mRNA analysis via branched DNA assay.

Quantigene 2.0 branched DNA Assay

Tissue punches were removed from RNA later and lysed with sterile beads in 300µL/sample of homogenizing solution (Quantigene sample processing kit, Invitrogen CAT#QG0517) and 6µL/sample of 20mg/mL proteinase K (Qiagen, mat #1114886). Lysis completed with a 30-minute incubation at 55°C. Lysates were removed from beads and stored at -80°C. On day of analysis, lysates were thawed and again incubated at 55°C for 30 minutes. Quantigene 2.0 probe sets prepared for human *HTT* (SA-50339) mRNA, mouse *Htt* (SB-14150) mRNA, and for mouse *HPRT* (SB-15463) mRNA as a housekeeping gene. 60µL of each probeset (mixture of mRNA specific probe, blocking reagent, Quantigene 2.0 lysis mixture, and water) was added to Quantigene plates (PSCP/HV DNA coated wells REF# 15553) in addition to 40µL of lysate. Quantigene plates were subsequently incubated overnight at 55°C.

On the following day, the plates were washed three times with wash buffer (Quantigene REF# 10843, and REF# 10846) and 100µL/well of pre-amplifier (Quantigene REF# 15095) was

added to the plate, which was then incubated for one hour at 55°C. The wash step was repeated and 100µL/well of amplifier (Quantigene REF# 15098) was added, and the 1-hour incubation was repeated. Following incubation, the plates were washed again, and 100µL/well of label probe (Quantigene REF# 13241) was added. Plates were then incubated for one hour at 50°C. The wash step was repeated one last time, and 100µL/well of substrate (Invitrogen REF# 144558) was added. Luminescence was measured within 15 minutes.

Background signal was subtracted, and all reads were normalized to PBS-injected control. Two-way ANOVA with Tukey's multiple comparisons test was performed using GraphPad Prism software. The significance is relative to the PBS injected control.

Western blot and WES

Frozen tissue punches were homogenized on ice in 75 µl 10mM HEPES pH7.2, 250mM sucrose, 1mM EDTA + protease inhibitor tablet (Roche) + 1mM NaF + 1mM Na3VO4, sonicated for 10 seconds and protein concentration determined by Bradford assay (BioRad). For western blot, equal amounts of protein (2.5 or 10 µg) were separated by SDS-PAGE on 4-12% Bis-Tris or 3-8% Tris acetate gels (Criterion XT, BioRad) and transferred to nitrocellulose using TransBlot Turbo (BioRad). Blots were probed with antibodies to HTT (aa1-17, Ab1⁷⁵), GFAP (Millipore), DARPP32 (abcam), Iba1 (Fujifilm) and reprobed with antibodies to loading controls Vinculin (Sigma) and GAPDH (Millipore). Bands were visualized with SuperSignal West Pico PLUS Chemiluminescent Substrate (Thermo) and Hyperfilm ECL (GE Healthcare). Total signal intensity was determined using the scanned films by manually tracing the bands in ImageJ software (NIH) and multiplying the area by the average signal intensity.

We also analyzed the same protein samples using the Wes system (ProteinSimple). The standard settings for the 66-440 kDa separation modules were used with 0.2 mg/ml of each lysate

and anti-HTT (Ab1, 1:50) plus anti-Vinculin (Sigma, 1:2000) antibodies. Peak area was determined using Compass for SW software (ProteinSimple) and dropped line fitted peaks.

Two-way ANOVA with Tukey's multiple comparisons test was performed on the normalized signal intensity and normalized peak area using GraphPad Prism software. Normalization and significance were calculated relative to the PBS injected control.

Thermo stability assay

Briefly, 1 μ M guide strand and 1 μ M complementary sense strand were annealed in a 10 mM sodium phosphate buffer (pH 7.2) containing 100 mM NaCl and 0.1 mM EDTA by heating at 95°C for 1 min and cooled down gradually to room temperature. Tm measurement was performed with temperature controller. Both the heating and cooling curves were measured over a temperature range from 20 to 95°C at 1.0°C/min three times⁶⁹.

Oligonucleotide synthesis

Oligonucleotides were synthesized by phosphoramidite solid-phase synthesis on a Dr Oligo 48 (Biolytic, Fremont, CA), or MerMade12 (Biosearch Technologies, Novato, CA), using 2'-F or 2'-OMe modified phosphoramidites with standard protecting groups. 5'-(E)-Vinyl tetra phosphonate (pivaloyloxymethyl) 2'-O-methyl-uridine 3'-CE phosphoramidite (VP) was purchased from Hongene Biotech, USA. All other phosphoramidites used were purchased from ChemGenes, Wilmington, MA. Phosphoramidites were prepared at 0.1 M in anhydrous acetonitrile (ACN), except for 2'-O-methyl-uridine phosphoramidite dissolved in anhydrous ACN containing 15% dimethylformamide. 5-(Benzylthio)-1H-tetrazole (BTT) was used as the activator at 0.25 M and the coupling time for all phosphoramidites was 4 min. Detritylations were performed using 3% trichloroacetic acid in dichloromethane. Capping reagents used were CAP A (20% n-methylimidazole in ACN) and CAP B (20% acetic anhydride and 30% 2,6-

lutidine in ACN). Reagents for capping and detritylation were purchased from American International Chemical LLC (AIC), Westborough, MA. Phosphite oxidation to convert to phosphate or phosphorothioate was performed with 0.05 M iodine in pyridine-H₂O (9:1, v/v) (AIC) or 0.1 M solution of 3-[(dimethylaminomethylene)amino]-3H-1,2,4-dithiazole-5-thione (DDTT) in pyridine (ChemGenes) for 4 min. Unconjugated oligonucleotides were synthesized on 500Å long-chain alkyl amine (LCAA) controlled pore glass (CPG) functionalized with Unylinker terminus (ChemGenes). Cholesterol conjugated oligonucleotides were synthesized on a 500Å LCAA-CPG support, where the cholesterol moiety is bound to tetra-ethylenglycol through a succinate linker (Chemgenes, Wilmington, MA). Divalent oligonucleotides (DIO) were synthesized on modified solid support¹⁹.

Deprotection and purification of oligonucleotides for screening of sequences

Prior to the deprotection, synthesis columns containing oligonucleotides were treated with 10% diethylamine (DEA) in ACN to deprotect cyanoethyl groups. In synthesis columns, both unconjugated and cholesterol conjugated oligonucleotides on solid support were then deprotected with monomethylamine gas (Specialty Gases Airgas) for an hour at room temperature. Deprotected oligonucleotides released from the solid support were precipitated on the support by passing solution of (i) a mixture of 0.1 M sodium acetate in 85% ethanol and then (ii) 85% ethanol to the synthesis column. The excess ethanol on solid support was dried by air flow and the oligonucleotides were flushed out by passing water through the column. This procedure renders pure oligonucleotides used for in vitro experiments.

Deprotection and purification of oligonucleotides for in vivo experiments

Prior to the deprotection, oligonucleotides on solid support were treated with 10% DEA in ACN in synthesis columns to deprotect cyanoethyl groups. Divalent oligonucleotides (DIO)

were cleaved and deprotected by AMA treatment for 2 hours at 45°C. The VP containing oligonucleotides were not treated with DEA post-synthesis and were cleaved and deprotected as described previously⁷⁶. Briefly, CPG with VP-oligonucleotides was treated with a solution of 3% DEA in 28-30% ammonium hydroxide at 35°C for 20 hours.

All solutions containing cleaved oligonucleotides were filtered to remove the CPG and dried under vacuum. The resulting pellets were re-suspended in 5% ACN in water. Purifications were performed on an Agilent 1290 Infinity II HPLC system. Crude oligonucleotides were purified using a custom 30x150mm column packed with Source 15Q anion exchange media (Cytiva, Marlborough, MA); running conditions: eluent A, 10 mM sodium acetate in 20% ACN in water; eluent B, 1 M sodium perchlorate in 20% ACN in water; linear gradient, 10 to 35% B in 40 min at 50°C. Flow was 30mL/min, peaks were monitored at 260 nm. Fractions were analyzed by liquid chromatography mass spectrometry (LC–MS). Pure fractions were combined and dried under vacuum. Oligonucleotides were re-suspended in 5% ACN and desalted by size exclusion on a 50×250 mm custom column, packed with Sephadex G-25 media (Cytiva, Marlborough, MA), and lyophilized.

LC–MS analysis of oligonucleotides

The identity of oligonucleotides was verified by LC–MS analysis on an Agilent 6530 accurate mass Q-TOF using the following conditions: buffer A: 100 mM 1,1,1,3,3,3-hexafluoroisopropanol (HFIP) and 9 mM triethylamine (TEA) in LC–MS grade water; buffer B: 100 mM HFIP and 9 mM TEA in LC–MS grade methanol; column, Agilent AdvanceBio oligonucleotides C18; linear gradient 0–35% B 5min was used for VP and DIO oligonucleotides); linear gradient 50–100% B 5min was used for TegChol conjugated oligonucleotides; temperature, 60°C; flow rate, 0.85 ml/min. LC peaks were monitored at

260nm. MS parameters: Source, electrospray ionization; ion polarity, negative mode; range, 100–3,200 m/z; scan rate, 2 spectra/s; capillary voltage, 4,000; fragmentor, 200 V; gas temp, 325°C

Deprotection, purification and LC-MS reagents were purchased from Fisher Scientific, Sigma Aldrich and Oakwood Chemicals.

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Conflict of Interest

FC, EP, NA and AK are inventors of a patent application related to SNP-based discrimination. AK and NA own stock in Atalanta Therapeutics. Other authors declare no conflict of interest.

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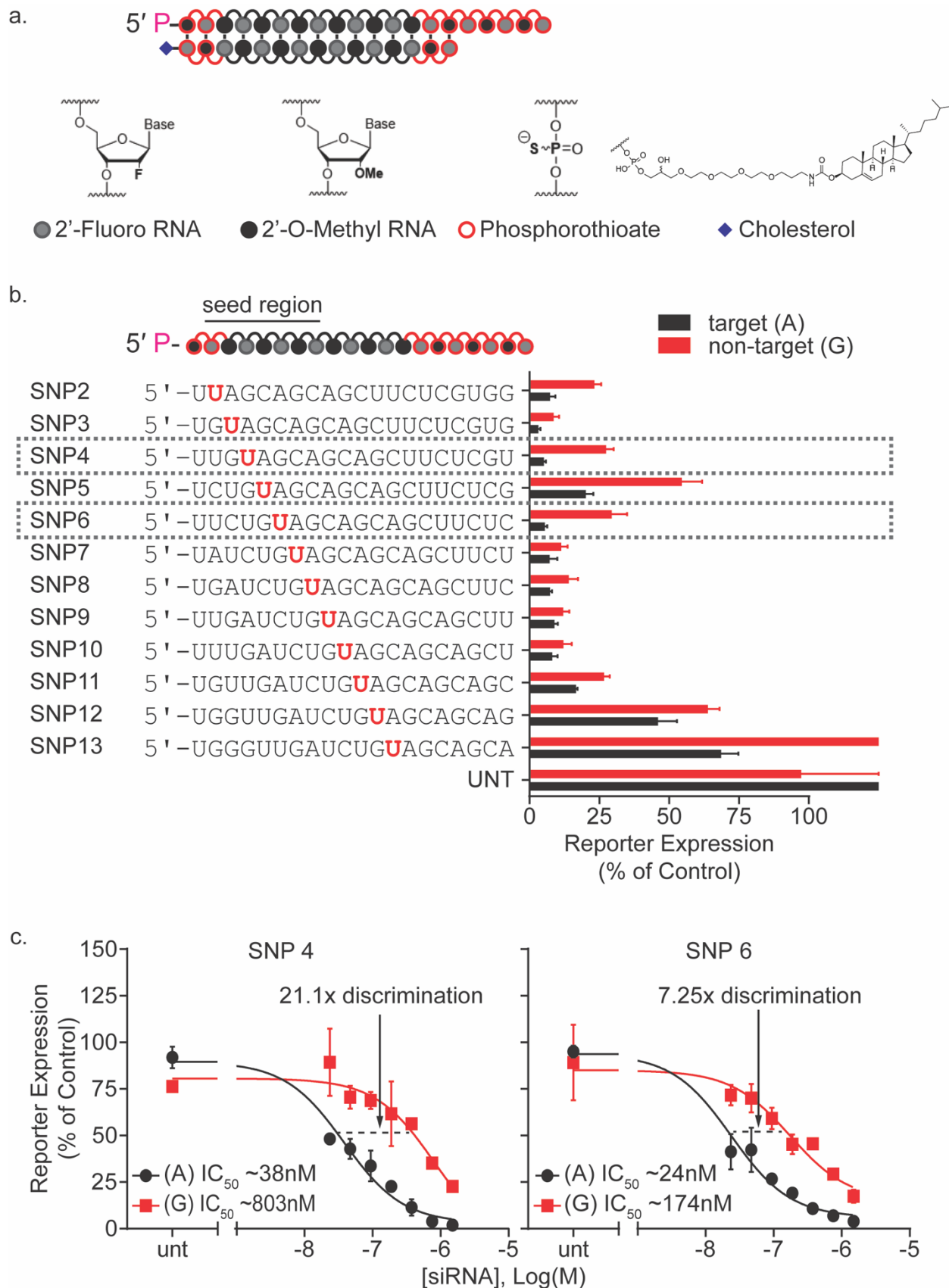


Figure 1. The primary screen for optimal siRNA sequence yields potent compounds with moderate discriminating power. (a) siRNA structure and chemical modification pattern used in screening (b) The primary screen identifies the most favorable positions of the SNP enabling single nucleotide discrimination for targeting of SNP site rs362273 (highlighted in red). Compounds were tested using a dual-luciferase reporter assay system in HeLa cells. The (psiCheck) reporter plasmids contain a 40mer region of huntingtin, including the target (A) SNP (black), and non-target (G) isoform (red). Cells were treated for 72 hours at 1.5 μ M of siRNA. A panel of siRNA sequences were synthesized in a cholesterol-conjugated scaffold with phosphorothioate and alternating 2'-F and 2'-OMe backbone modifications. By walking the siRNA sequence around SNP site rs362273, we find multiple compounds with varying degrees of efficacy and discrimination. (c) 7-point dose-response shows that siRNAs with the SNP site in positions 4 (SNP4) and 6 (SNP6) generate 20-fold and 7-fold allelic discrimination, respectively, with a high degree of efficacy.

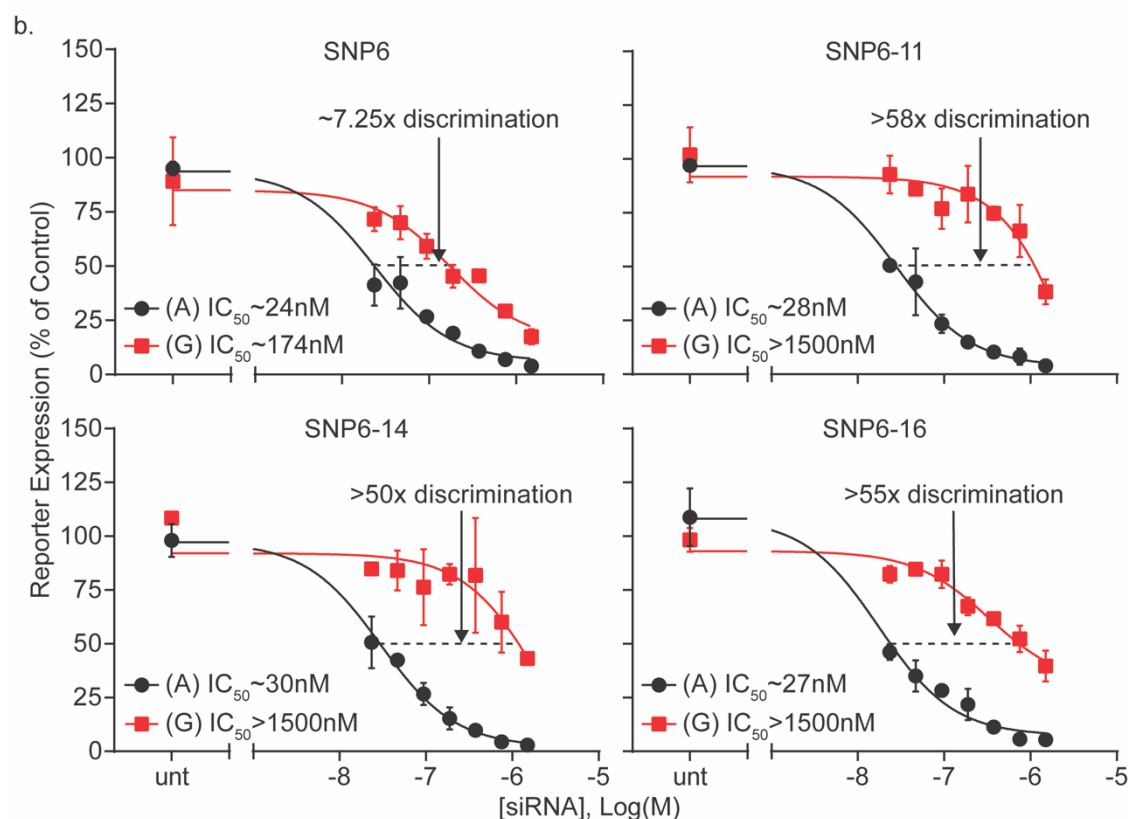
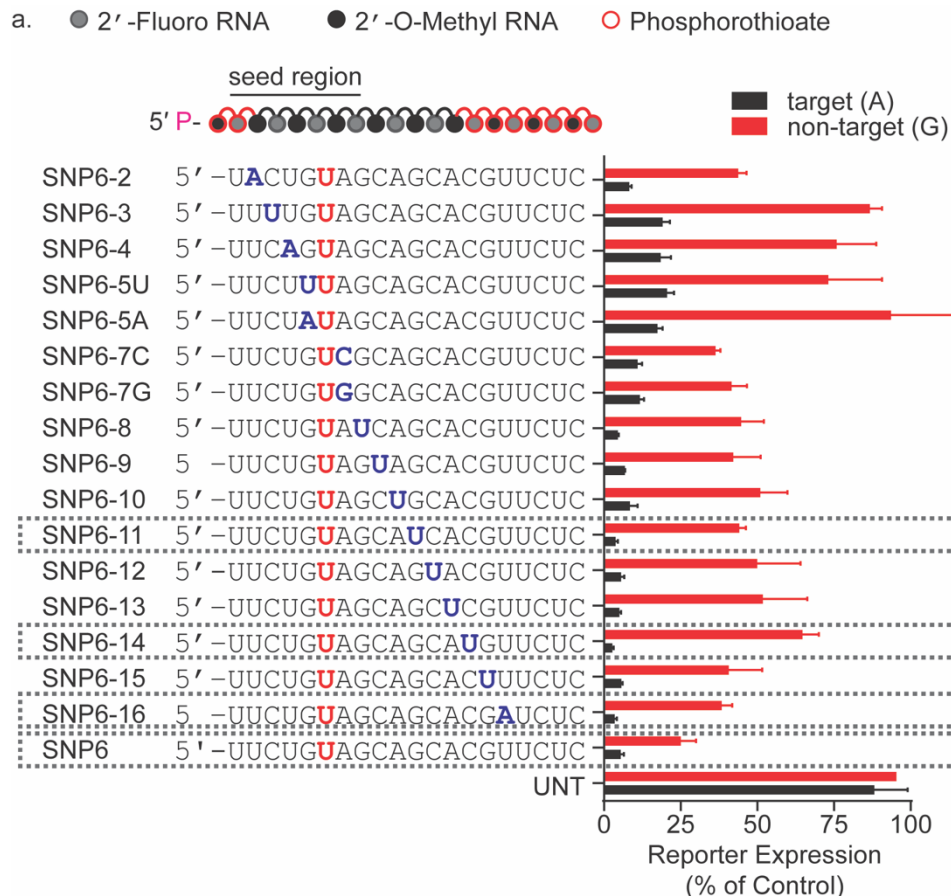


Figure 2. A systematic screen of siRNAs targeting a heterozygosity at site rs362273 yields multiple compounds in which a secondary mismatch improves allelic discrimination. (a) In certain positions, adding an intentional mismatch to a SNP-targeting siRNA improves discrimination without impairing target silencing. Compounds were tested using a dual-luciferase reporter assay system in HeLa cells. The (psiCheck) reporter plasmids contain a 40mer region of huntingtin, including the target (A) SNP isoform, and non-target (G) SNP isoform. Cells were treated for 72 hours at 1.5 μ M of siRNA. An siRNA selected from the primary screen (SNP6) (Fig.1) was used as a template sequence and a secondary mismatch (highlighted in blue) was introduced. When a mismatch is added outside the seed region, siRNAs show a substantial increase in discriminating power without decreasing target silencing. (b) 7-point dose-response of selected compounds shows that the addition of a U:C mismatch in position 11 increases discrimination from approximately 7-fold to greater than 50-fold. Changes in efficacy and discrimination vary based on the position of the secondary mismatch.

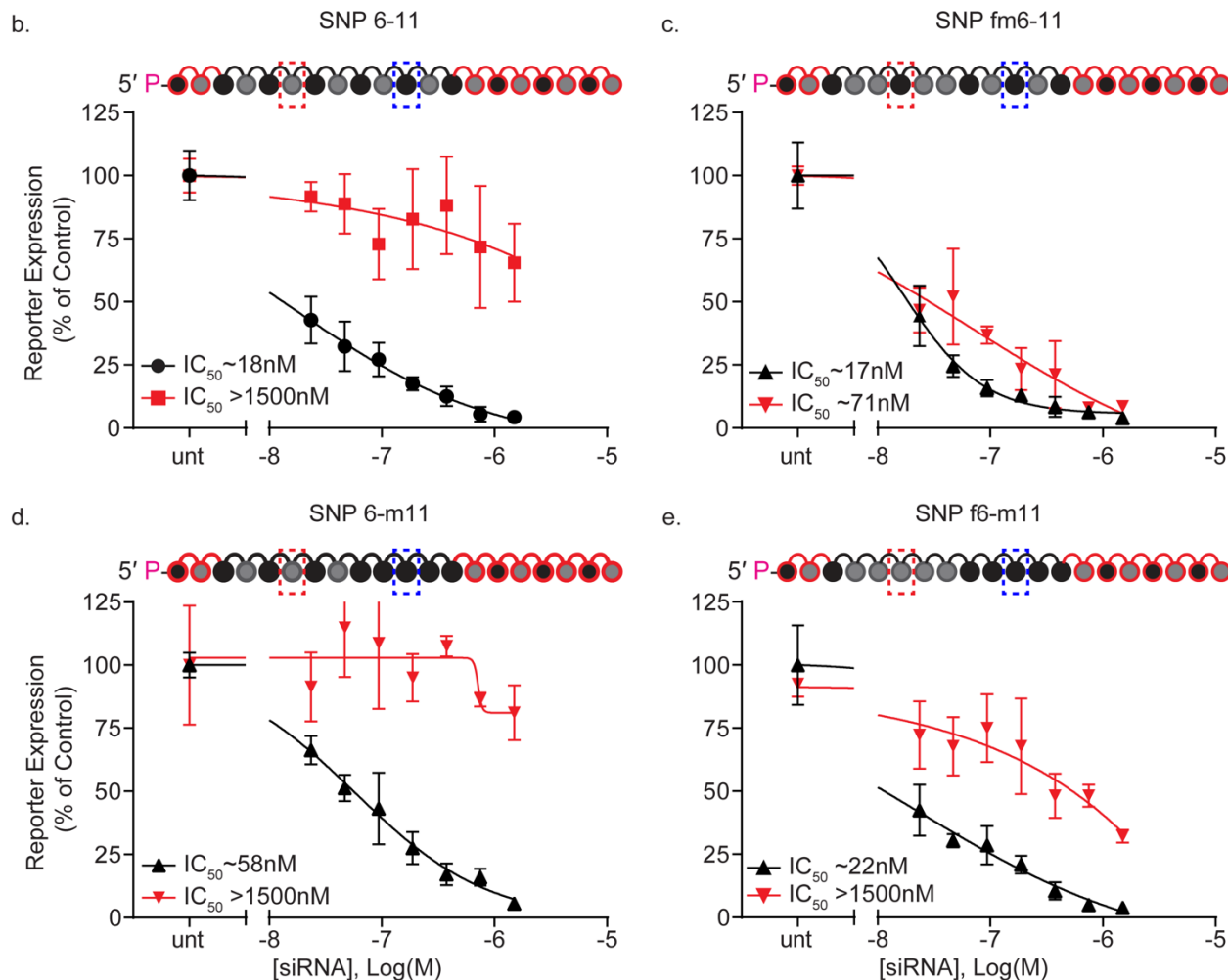
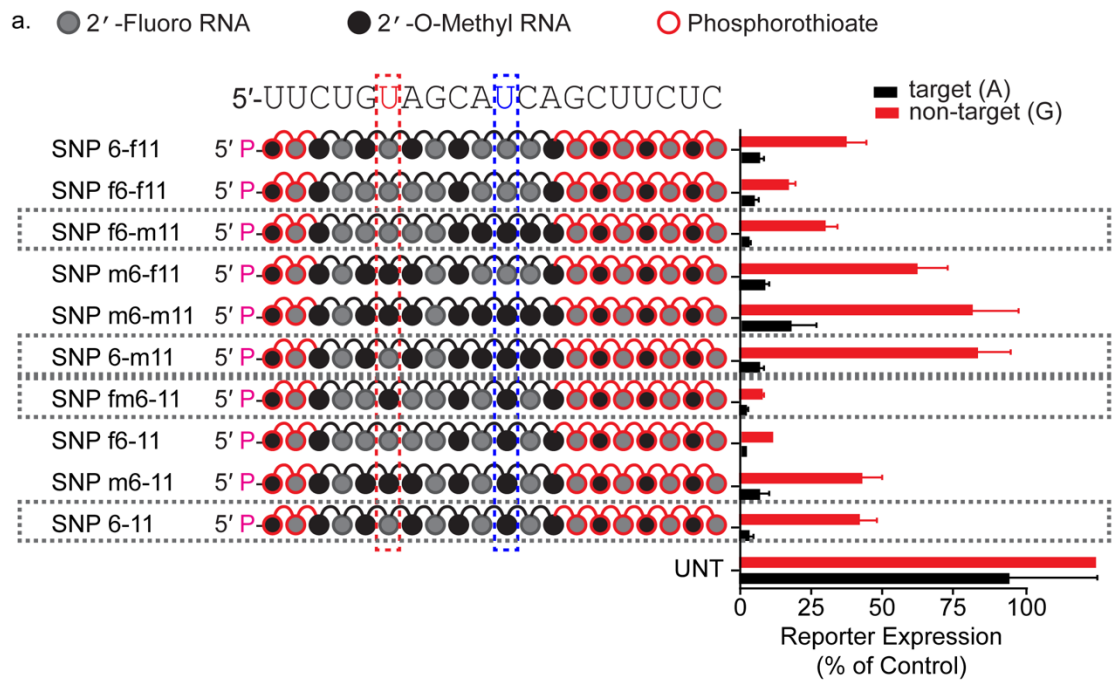


Figure 3. Chemical modification pattern affects allele specificity and efficacy. (a)

Combinations of 2'-F and 2'-OMe enrichment around critical positions 6 and 11 affect siRNA activity. Compounds were tested using a dual-luciferase reporter assay system in HeLa cells. The (psiCheck) reporter plasmids contain a 40mer region of huntingtin, including the target (A) SNP isoform, and non-target (G) SNP isoform. Cells were treated for 72 hours with 1.5 μ M siRNA. **(b-e)** 7-point dose-response of alternative modification patterns show that backbone modification pattern has impacts efficacy and discrimination. Compared to the original alternating modification pattern **(b)**, a heavily-fluorinated SNP region shows an increase in target efficacy, but a complete loss of discrimination between alleles **(c)**. A heavily methylated mismatch region increases discrimination but decreases target efficacy **(d)**. Combining a fluorinated SNP region with a methylated mismatch region improves efficacy but does not increase discrimination above the original alternating modification pattern **(e)**.

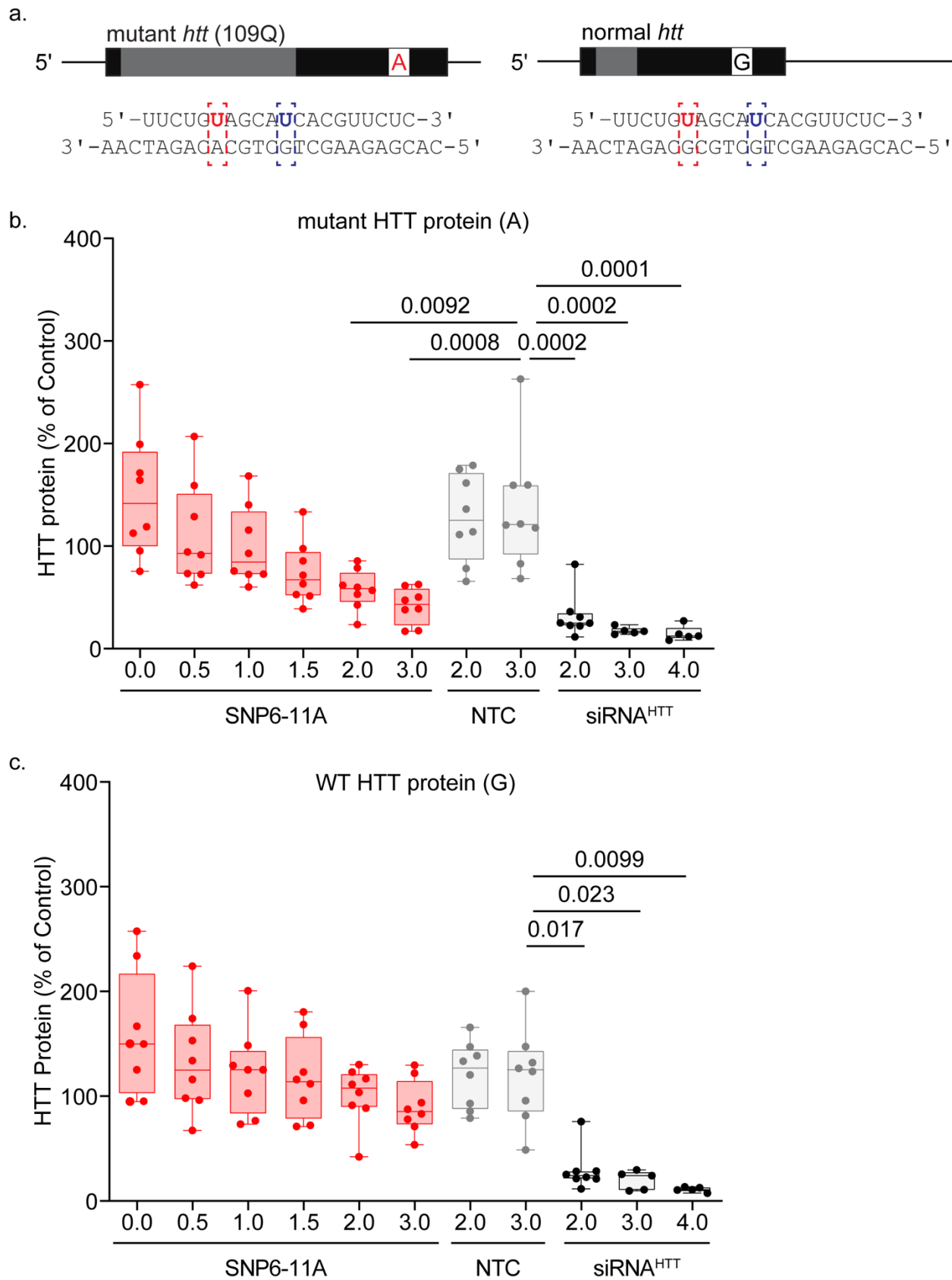


Figure 4. SNP6-11A siRNA maintains selective-silencing of mutant HTT in patient-derived NSCs HD patient derived neural stem cells carrying an A or a G at SNP site rs362273 (a) were treated for 5 days with SNP6-11 or non-selective siRNA^{HTT} at the concentrations listed. (b,c) Graphs show pixel intensity quantification of mutant (b), or wild-type (c) HTT signal intensity standardized to that of vinculin reported as a percentage of untreated NSCs. A non-targeting control (NTC) siRNA was used as a negative control. Non-selective targeting with siRNA^{HTT} was used as a positive control. ANOVA and posthoc Tukey test. Error bars, SD; N=5-8.

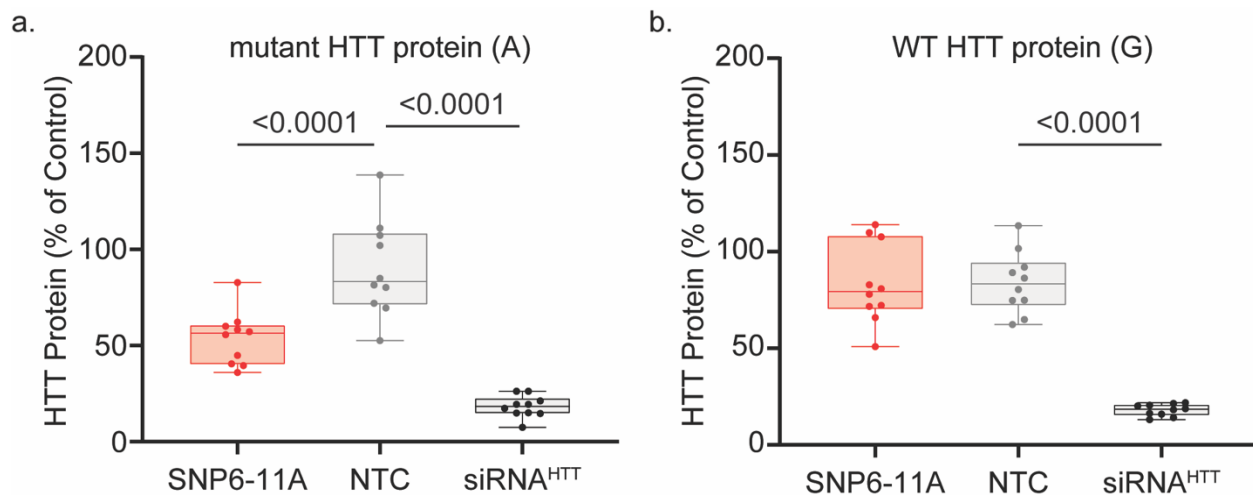


Figure 5. SNP6-11A siRNA maintains selective-silencing of mutant HTT in neurons differentiated from human iPSCs. Western blot analysis of HD109 neurons, which are heterozygous at SNP2273, showed lowering of mutant HTT (a) compared to wild-type HTT (b) with SNP6-11. Cells were treated for 5 days with 3.0 μ M siRNA. Pixel intensity of mutant or wild-type HTT was standardized to that of vinculin and reported as a percentage of untreated. Non-selective targeting with siRNA^{HTT} was used as a positive control. A non-targeting control (NTC) siRNA was used as a negative control. ANOVA and posthoc Tukey test, n=10.

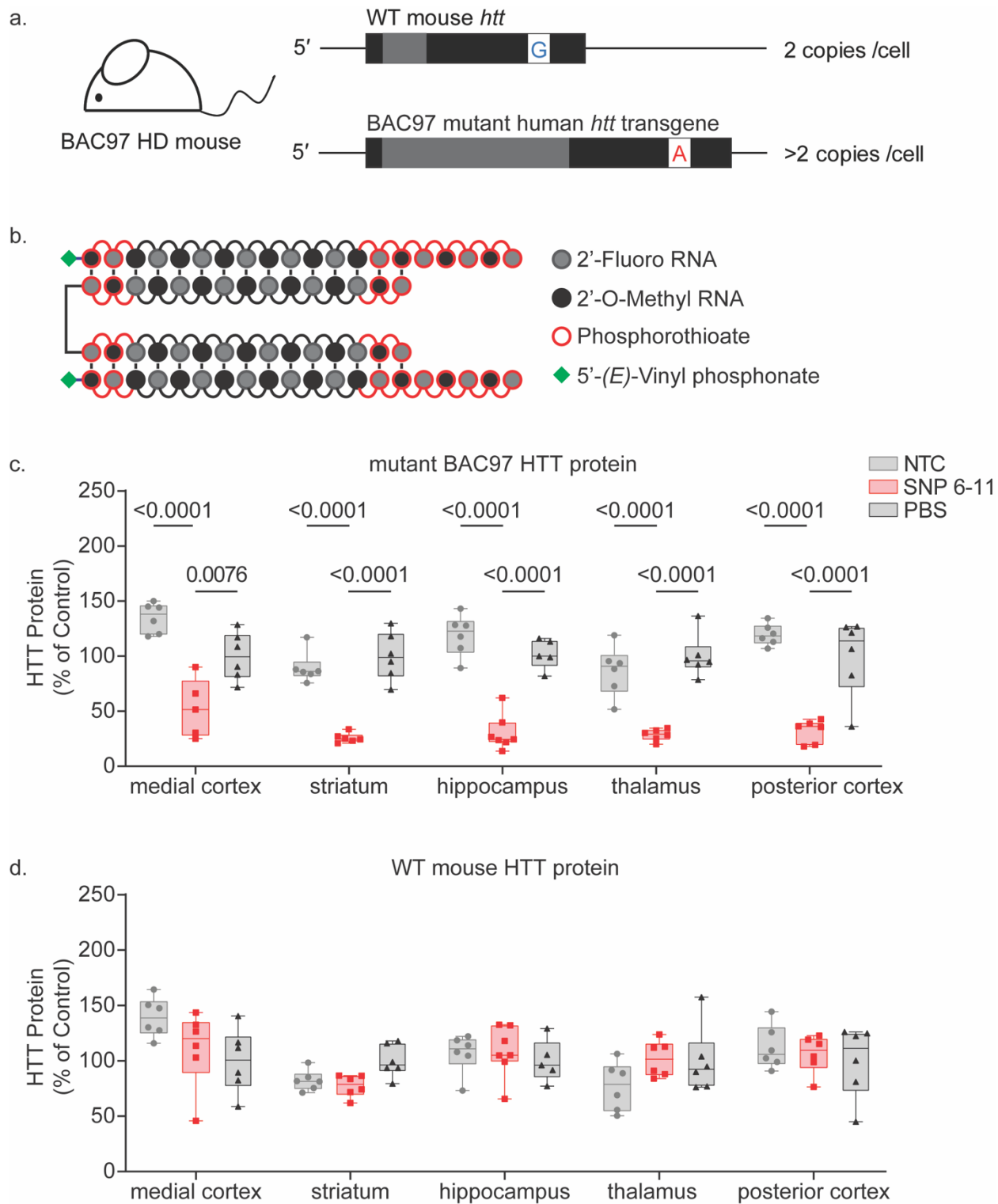


Figure 6. SNP6-11 selectively silences mutant HTT in an HD mouse model. (a) BAC97 HD mice have a high copy number of transgenic mutant human huntingtin (*Htt*) with a polyQ expansion of 97 repeats, and two normal copies of wild-type (WT) mouse *Htt*. Both the human BAC97 huntingtin and the WT mouse *Htt* have homology at the SNP6-11 target site, except for a heterozygosity at SNP site rs362273, mimicking the WT/mutant *Htt* SNP heterozygosity in

patients. **(b)** for *in vivo* delivery, the SNP 6-11 compound was synthesized in a di-valent configuration as depicted **(c)** When treated with 225 μ g (10nmols) of siRNA (SNP6-11) in a divalent scaffold, allele-specific silencing of mutant HTT protein in mouse brain is achieved one month after bilateral injection into the lateral ventricles (ICV injection). **(d)** Across brain regions, WT mouse HTT is preserved with no significant silencing detected (except in striatum), while transgenic mutant HTT is lowered significantly, averaging 70% silencing across brain regions (p-value <0.00001). Protein levels were measured via WES Protein Simple. A two-way ANOVA with multiple comparisons was performed for all protein analysis, comparing treatment groups to the PBS control for each brain region.