Development of a Clinically Viable Heroin Vaccine

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Supporting Information

ABSTRACT: Heroin is a highly abused opioid and incurs a significant detriment to society worldwide. In an effort to expand the limited pharmacotherapy options for opioid use disorders, a heroin conjugate vaccine was developed through comprehensive evaluation of hapten structure, carrier protein, adjuvant and dosing. Immunization of mice with an optimized heroin-tetanus toxoid (TT) conjugate formulated with adjuvants alum and CpG oligodeoxynucleotide (ODN) generated heroin “immunoantagonism,” reducing heroin potency by >15-fold. Moreover, the vaccine effects proved to be durable, persisting for over eight months. The lead vaccine was effective in rhesus monkeys, generating significant and sustained antidrug IgG titers in each subject. Characterization of both mouse and monkey antiheroine antibodies by surface plasmon resonance (SPR) revealed low nanomolar antiserum affinity for the key heroin metabolite, 6-acetylmorphine (6AM), with minimal cross reactivity to clinically used opioids. Following a series of heroin challenges over six months in vaccinated monkeys, drug-sequestering antibodies caused marked attenuation of heroin potency (>4-fold) in a schedule-controlled responding (SCR) behavioral assay. Overall, these preclinical results provide an empirical foundation supporting the further evaluation and potential clinical utility of an effective heroin vaccine in treating opioid use disorders.

INTRODUCTION

Heroin, a semisynthetic opioid, and its parent natural product, morphine, are among the longest known and commonly abused psychoactive drugs. Heroin is a prodrug1 that readily crosses the blood-brain barrier while quickly deacetylating to 6-acetyl morphine (6AM) and then more slowly to morphine (Figure 1).2,3 These two metabolites agonize brain mu-opioid receptors (MORs) to produce heroin’s abuse-related euphoric and reinforcing effects.4−6 Moreover, the robust analgesic effects of opioids have led to their extensive clinical use as prescription painkillers such as OxyContin (oxycodone) and Vicodin (hydrocodone); however, these opioids are also routinely abused and can act as “gateway drugs” to heroin.7,8 Persistent opioid abuse leads to a neuropsychiatric disorder, i.e., opioid use disorder, characterized by compulsive opioid administration despite the negative physical, mental, legal and social consequences of prolonged use.

Currently in the United States, opioid abuse has reached epidemic levels. The number of people who have used heroin in the past 10 years has doubled from 379,000 in 2005 to 828,000 in 20159,10 and heroin expenditures have grown steadily to an estimated $27 billion (2010) on drug purchases alone.11 The widespread prevalence of heroin abuse is a significant cost to users and to society as a whole (an estimated total of $22 billion in the US).12,13 Other negative impacts of heroin abuse include HIV or HCV infection, for which injection drug users remain at the highest risk.14 While prescription opioids combined are involved in the most drug-related deaths in the US, compared to any one single drug, heroin is responsible for at least twice as many deaths.15 Abuse of prescription opioids may be mitigated by tightening regulations or by introducing antiabuse technology during manufacturing. On the other hand, heroin and other synthetic opioids, e.g., acetyl fentanyl, are produced and distributed illegally; therefore, great measures must be taken to curb illicit opioid use. Current treatment options for opioid use disorders include opioid replacement therapy utilizing methadone or buprenorphine as MOR agonists to reduce opioid withdrawal symptoms and maintain heroin abstinence.16,17 Opioid antagonists naltroxone and naltrexone (NTX) are other treatment options, FDA-approved for opioid overdose and dependence, respectively.18,19 Pharmacological intervention for heroin abuse has proven to be effective but has a number of drawbacks including high cost of in-patient rehab.20,21
undesirable effects, and relapse potential following therapy. 

Humanity has benefited from vaccines for more than two centuries, and of all the biomedical achievements, immunization for the prevention of infectious diseases ranks highly. The first attempt at translating vaccination to reduce the use of psychoactive substances was reported in the early 70s when a conjugate vaccine containing a morphine-like hapten was tested in a single rhesus monkey. However, this work was not followed up due to the emergence of pharmacotherapies for opioid use disorders, e.g., methadone, which at the time, appeared more promising. Drug conjugate vaccine research re-emerged in the mid-90s, and focused on cocaine and nicotine. Unfortunately, multiple failures of both cocaine and nicotine vaccines in human trials have called into question the clinical value of vaccination for treating substance use disorders. Potential problems of these vaccines include poor hapten design and adjuvant selection. Moreover, these vaccines lacked rigorous preclinical development, as they have not demonstrated the ability to block a wide range of drug doses in multiple behavioral procedures. Failure to address and ultimately solve these problems has hampered progress in the drug-vaccine field.

The principle design elements behind drug vaccines include a hapten (B-cell epitope), highly congruent in structure to the target drug, and an immunogenic carrier protein (T-cell epitope) such as tetanus toxoid (TT). Immunization of the hapten-protein conjugate formulated with adjuvants, e.g., alum and CpG oligodeoxynucleotide (ODN), triggers an adaptive immune response against the drug-like hapten. Subsequently, when the vaccinated subject receives a drug dose, available polyclonal IgG antibodies in the periphery bind the drug with a high degree of affinity and specificity, precluding drug entry to the brain. The unique mechanism of action of drug-conjugate vaccines may offer many advantages for treating substance use disorders such as the potential for reduced side effects, convenient and low-cost administration, and long-term efficacy.

We envisioned the need for a comprehensive series of preclinical experiments wherein every component of the vaccine would be scrutinized, i.e., hapten, carrier and adjuvant. Moreover, the cornerstone of our development process involved the quantitative evaluation of each vaccine iteration in a rodent antinociception assay. Thus, full heroin dose–response curves were generated to compare ED50 values of vaccinated and nonvaccinated groups of rodents, providing a direct measure of the “immunoantagonistic” capacity of the vaccine. A first-generation heroin vaccine produced heroin ED50 ratios of 4–5 in both mice and rats, blocking the effects of heroin in a series of behavioral models. In this work, a significant leap forward was made in redesigning the vaccine to achieve a greater than 15-fold heroin ED50 shift in rodents, warranting further investigation in a primate behavioral model. The translational potential of the heroin vaccine was confirmed for the first time in rhesus monkeys, supported by statistical analysis in n = 4 subjects.
RESULTS

Vaccine Optimization: Hapten, Carrier and Adjuvant.

As depicted in Scheme 1, preparation of heroin hapten was accomplished by first demethylating heroin via Olofson’s procedure. Reductive amination of Boc-protected 4-amino-butanal followed by TFA deprotection afforded the key intermediate 1 as previously described. Amidation coupling(s) followed by trityl or t-butyl ester deprotections yielded the first generation thiol hapten \( (\text{SH}) \) or novel second generation carboxylic acid hapten (2-4). Thiol and carboxylic acid hapten were coupled to surface lysines of carrier proteins via thiol-maleimide or amide couplings, respectively. At a 1:1 w/w ratio of hapten to protein, carboxylate hapten showed higher hapten loading according to MALDI-ToF analysis (Table S1, Figure S1). Additional advantages to carboxylate hapten conjugations included resistance to oxidation and a one-pot coupling procedure. In contrast, thiols can form disulfides and require preparation of maleimide-loaded proteins prior to conjugation, i.e., a two-pot procedure.

Figure 3. Optimization of heroin immunoconjugate and vaccine formulation. (A) Comparison of carrier proteins with HerSH hapten. \(^{**}P < 0.01, ^{*}P < 0.001, ^{**}P < 0.001\) versus KLH. (B) Evaluation of heroin hapten as TT immunoconjugates. \(^{***}P < 0.001, ^{***}P < 0.001\) versus HerSH-TT. (C) Comparison study of 6AMCOOH vs HerCOOH hapten as TT conjugates. \(^{**}P < 0.01, ^{***}P < 0.001\) versus HerSH-TT. (D) Dose-dependency of HerCOOH-TT immunoconjugate on vaccine efficacy. \(^{**}P < 0.01, ^{***}P < 0.001, ^{**}P < 0.05\) versus 10 μg dose. No CpG ODN was used. (E) Adjuvant effects of CpG ODN 1826 and cGAMP on vaccine efficacy. Doses: 4 μg CpG (Low CpG), 30 μg CpG (High CpG), 60 μg CpG + 100 μg HerCOOH-TT (2X dose), 1.3 μg (Low cGAMP), 10 μg (High cGAMP). \(^{*}P < 0.05, ^{**}P < 0.01, ^{***}P < 0.01, ^{***}P < 0.001\) versus alum only. Vaccine formulations for all panels contained 0.75 mg alum, 30 μg CpG ODN 1826 and 50 μg HerCOOH-TT unless otherwise noted and were administered to \( n = 6 \) mice i.p. at wk 0, 2, and 4. For all panels, the mean heroin ED\(_{50}\) ± SEM (determined via cumulative s.c. dosing) is shown and statistics were evaluated by a one-way ANOVA with Tukey’s posthoc test. Testing was performed at wk 6 and significance is denoted by \(^{(*)}\) for tail immersion and \(^{(\#)}\) for hot plate. Fold-shifts in ED\(_{50}\) versus \( n = 6 \) nonvaccinated control mice are reported above each set of data points. Raw antinociception curves shown in Figures 3A-E.
compared through the testing of TT conjugates of HerSH (5), HerCOOH (2) and HerdBA (4), and the HerCOOH hapten demonstrated the greatest efficacy (Figure S2B). While the amide coupling method appeared to generate a more efficacious conjugate compared to thiol-maleimide coupling, this benefit appeared to be erased by the presence of the di-beta-alanine (dBA) linker found in the HerdBA hapten. The effect of the 3-acetyl group was probed through a 6AM-like hapten (6AMCOOH, 3), because 6AM is the main mediator of heroin psychoactivity.4 Behavioral results indicated that the 6AM and the heroin haptenens were comparable in efficacy (Figure 3C). ELISA results corroborated the behavior because sera from both groups bound both 6AM and heroin haptenens to a similar degree; however, the 6AM conjugate elicited antibodies with a slightly reduced capacity to bind the heroin hapten (Figure S2C).

In continuing studies with the HerCOOH-TT conjugate, the effect of conjugate dosing and adjuvant was explored, using FDA-approved alum adjuvant (Al(OH)₃) alone as a benchmark. Conjugate dosing demonstrated a positive effect on titer levels and mitigation of heroin antinociception (Figure S2D, 3D). Addition of CpG ODN 1826, a Toll-like receptor (TLR) 9 agonist, enhanced vaccine potency of HerCOOH-TT (Figure 3E) as previously shown for HerSH-KLH.45 Without alum, CpG ODN was not effective (data not shown). When both conjugate and CpG ODN doses were doubled to 100 µg and 60 µg respectively, heroin “immunoantagonistic” capacity and antihapten titer levels increased dramatically (Figure 3E, S2E). A second DNA-based immunostimulatory adjuvant, 2′3’-cyclic guanosine monophosphate-adenosine monophosphate (cGAMP)49 was also evaluated. Although both cGAMP and CpG ODN target the innate immune system, they activate different receptors and pathways via STING and TLR9, respectively. A mild vaccine improvement was noted with low dose cGAMP, however, this effect was not present at a higher cGAMP dose (Figure 3E). The outcome of these optimization studies was the identification of a lead vaccine formulation containing HerCOOH-TT, CpG ODN and alum.

**Long-Term Duration of Lead Vaccine Efficacy.** In order to test the durability of the optimized heroin vaccine, an extended vaccination study was performed over a 37-week period in mice. After an initial immunization, peak antihapten titers were observed by ELISA, causing significant potency shifts in heroin ED₅₀ as measured by antinociceptive testing (Figure 4A–C). Although vaccine-mediated shifts in heroin ED₅₀ declined over the next three months, a second and third round of vaccinations at months 5 and 8 maintained vaccine efficacy at approximately 50–70% of the initial level (Figure 4A–C). As a means to corroborate the antinociception results, ELISA titers were evaluated with the caveat that serum binding to an immobilized drug hapten does not necessarily equate to affinity for the actual drug molecule.40–42 To address this potential limitation, individual heroin ED₅₀s were plotted against antihapten titers, and a linear relationship was observed (Figure 4D). The correlation implies that the degree of serum antibody binding to the HerCOOH hapten is representative of the degree of binding to the actual opioids (heroin and 6AM) in vivo.

A possible liability in using TT in the final formulation is that preexisting immunity to TT from the clinically administered DTaP vaccine could reduce the subsequent response to a heroin vaccine, a phenomenon known as carrier-induced epitopic suppression (CIES).43,44 However, no evidence was found for the occurrence of CIES in the context of the current heroin vaccine (Figure S4).
Figure 5. Heroin conjugate vaccine elicits a robust antidrug antibody response in monkeys with high affinity and selectivity for 6AM. (A) Binding curves of vaccinated monkey and mouse antisera for heroin and 6AM as determined by SPR in triplicate. Points represent the mean binding to immobilized HerCOOH-BSA following serum incubation with 12 dilutions of heroin or 6AM competitors. Binding values were normalized to serum binding without competitor drug. Listed IC_{50} ± SEM were derived from a nonlinear fit of the binding curves. Serum was pooled from n = 2 monkeys (M1,2) and n = 6 mice collected after three initial immunizations. (B) Binding selectivity of mouse and monkey antisera for various opioids at 100 μM ([6AM] = 10 μM) by SPR in triplicate. Minimum selectivity factors relative to 6AM are shown above each bar. (C) Antiteroid IgG titers of rhesus monkeys (M1,2) vaccinated at the indicated time points (arrows) with 400 μg HerCOOH-TT, 600 μg CpG ODN 2006 and 5 mg alum. Midpoint titers were determined in duplicate by ELISA against a HerCOOH-BSA coating antigen. No antiheroin IgM titers were detected at any point. Monkeys M1 and M2 were previously vaccinated with the heroin vaccine in a pilot study while M3 received unconjugated TT. M4 was not pretreated. (D) Individual monkey antisera affinity for 6AM over time by SPR. Points represent mean IC_{50} ± SEM determined from a 12 point 6AM dilution curve similar to panel A methods. Time points match up with the panel C timeline.

**Heroin Vaccine-Induced Antibody Response in Non-human Primates (NHPs).** As an initial assessment of the lead vaccine in NHPs, a pilot vaccine study was performed in which two rhesus monkeys (M1,2) received the heroin conjugate while one monkey (M3) received the unmodified TT carrier protein as a control. Following three injections, a significant and consistent antidrug IgG antibody response was observed in the conjugate-vaccinated monkeys while antheroin antibodies were not observed in the carrier-vaccinated monkey (Figure S5A). Preliminary behavioral assessments suggested that heroin dose-effect curves were right-shifted >3-fold from baseline in both conjugate-vaccinated monkeys (Figure S5B). Surface plasmon resonance (SPR) analysis of both monkey and mouse sera indicated submicromolar and low nanomolar competitive IC_{50} for heroin and 6AM, respectively (Figure S5A), which in contrast to ELISA is representative of actual antibody K_{A}. In comparing binding selectivity for other opioids, antisera affinity for morphine, oxycodone and methadone was >1000-fold lower (Figure S5B).

An extended vaccination study involving the same monkeys including one new one (M4) demonstrated that all subjects produced a long-lasting, high IgG titer response to the vaccine (Figure 5C). Interestingly, the two monkeys that received the heroin vaccine in the pilot study (M1,2) showed significantly higher titers than the monkeys that received carrier (M3) or no vaccine (M4). SPR analysis of antisera from each monkey revealed consistent drug affinity for 6AM of ≤1 nM in monkeys M1,2 while monkeys M3,4 gradually increased 6AM affinity over the course of the study to ~6 nM (Figure 5D). The 6AM affinity of M3,4 matches the 6AM affinity of M1,2 observed after the pilot study (Figure S5A), while 6AM affinity in M1,2 increased further by approximately 6-fold following the second immunization study (Figure 5D).

**Vaccine-Mediated Alteration of Heroin Pharmacology in NHPs.** Given the strong presence of 6AM-neutralizing antibodies in vaccinated rhesus monkeys, further experiments were conducted to evaluate the capacity of the antibodies to alter the pharmacodynamics (PD) and pharmacokinetics (PK) of heroin. MOR agonists consistently produce dose-dependent decreases in operant responding that appear to be mediated by pharmacologically similar populations of MORs. These MORs mediate other MOR agonist behavioral effects, such as antinociceptive and discriminative stimulus effects. Moreover, drug ED_{50} values can be quantified from operant responding to serve as a potency metric for MOR modulators such as the antagonist NTX; therefore, an assay of schedule-controlled responding (SCR) was selected as a reliable behavioral measure of opioid pharmacology upon which to examine the “immunoantagonist” vaccine.

Following the double-determination of baseline heroin and oxycodone potencies to decrease rates of responding (represented by ED_{50} values), monkeys were vaccinated and retested 6 weeks later, resulting in a clear 3.5-fold heroin ED_{50} shift (Figure 6A). A greater potency ratio was detected 2–3 weeks following week 11 (4.3-fold) and 18 (41-fold) booster injections (Figure 6A, SSC). The heroin ED_{50} values at these times (weeks 14 and 21) were similar to heroin ED_{50} values obtained following acute NTX pretreatment (Figure 6B). Not only were heroin potency ratios significantly increased versus baseline levels, they were also selectively elevated compared to the control opioid, oxycodone, throughout the entire study (Figure 6A, S6). A correlation between monkey antitheroin
Correlation between titers against HerCOOH-BSA and SCR heroin ED\textsubscript{50} values collected over 26 weeks. Comparison to naltrexone (NTX) treatment at 3.2 μg/kg (1x) and 0.19 ± 0.06 mg/kg, respectively. RM one-way ANOVA of heroin and oxycodone SCR over time: F\textsubscript{1,2},14 = 2.76 and 1.07, P = 0.0092 and 0.414, respectively; half-filled (P < 0.05) and fully filled (P < 0.01) circles indicate significance by Dunnett’s post hoc test vs heroin baseline. RM two-way ANOVA of heroin versus oxycodone SCR over time: F\textsubscript{1,3} = 13.2, P = 0.0038; half-filled (P < 0.05) and fully filled (P < 0.01) squares indicate significance by Bonferroni’s post hoc test versus heroin. (B) SCR cumulative heroin dose–effect curves at week 0 (baseline) and 14 (vaccine) in comparison to naltrexone (NTX) treatment at 3.2 μg/kg (1x) and 32 μg/kg (10x). Points represent mean ± SEM for n = 4 monkeys. (C) Correlation between titers against HerCOOH-BSA and SCR heroin ED\textsubscript{50} values collected over 26 weeks. P < 0.0001 by Pearson’s correlation. (D) 6AM serum concentrations over time in n = 4 rhesus macaques (M1–4) following 0.32 mg/kg i.m. heroin. The PK study was performed in the same subjects before and after a course of three initial immunizations. F\textsubscript{1,2,8} = 20.35, P = 0.0041 by a RM two-way ANOVA comparing pre and post vaccination PK (post-vacc 1). After 7 months, monkeys M1 and 2 received three additional immunizations and the PK study was repeated (post-vacc 2). (E) 6AM AUC values corresponding to the PK study. Fold increases versus baseline are reported as mean ± SEM. **P < 0.01 by paired t-test. C\textsubscript{max} values (ng/mL): baseline = 32.5 ± 6.8, post-vacc 1 = 418 ± 110, post-vacc 2 = 5430 ± 2800.

**DISCUSSION**

Achievement of heroin vaccine effectiveness required mastery of a number of unique challenges. First, the desired immune response from the heroin vaccine is distinct compared to vaccines being developed against pathogens because drug vaccines have a greater requirement for strong humoral immunity; a larger molar quantity of drug relative to pathogens must be neutralized by IgGs for the drug vaccine to achieve efficacy. On the other hand, while cell-mediated immunity is necessary for subduing disease-causing pathogens, it would be counterproductive in the development of immunoantagonists for the treatment of substance use disorders. Second, utilizing TT as a carrier protein at the clinical level could readily be questioned if pre-existing immunity to TT diminished the immune response to the drug-carrier conjugate. Fortunately, pre-existing antibody titers to TT did not suppress heroin vaccine efficacy. Third, as heroin is a prodrug for 6AM/morphine, hapten design was tailored to direct antibody binding toward 6AM followed by heroin but not morphine. Despite being a psychoactive metabolite of heroin, morphine penetrates the blood-brain barrier much less readily, hence, antibody sequestration of heroin/6AM until enzymatic hydrolysis ensues is key to vaccine performance.\textsuperscript{51} Fourth,

**Figure 6.** Heroin vaccine diminishes heroin potency and alters 6AM pharmacokinetics in rhesus monkeys. (A) Timeline of changes in either heroin or oxycodone ED\textsubscript{50} in monkeys after heroin vaccine administration. Monkeys (M1–4) were vaccinated i.m. at the indicated time points (arrows) with 400 μg HerCOOH-TT, 600 μg CpG ODN 2006 and 5 mg alum. Baseline heroin and oxycodone ED\textsubscript{50} values were double-determined in each monkey prior to vaccination in the assay of schedule-controlled responding (SCR). Each point in the figure represents the average ratio of heroin (red) or oxy (blue) ED\textsubscript{50} ± SEM relative to the baseline value. Group mean ± SEM baseline ED\textsubscript{50} values for heroin and oxycodone were 0.08 ± 0.03 and 0.19 ± 0.06 mg/kg, respectively. RM one-way ANOVA of heroin and oxycodone SCR over time: F\textsubscript{1,2},14 = 2.76 and 1.07, P = 0.0092 and 0.414, respectively; half-filled (P < 0.05) and fully filled (P < 0.01) circles indicate significance by Dunnett’s post hoc test vs heroin baseline. RM two-way ANOVA of heroin versus oxycodone SCR over time: F\textsubscript{1,3} = 13.2, P = 0.0038; half-filled (P < 0.05) and fully filled (P < 0.01) squares indicate significance by Bonferroni’s post hoc test versus heroin. (B) SCR cumulative heroin dose–effect curves at week 0 (baseline) and 14 (vaccine) in comparison to naltrexone (NTX) treatment at 3.2 μg/kg (1x) and 32 μg/kg (10x). Points represent mean ± SEM for n = 4 monkeys. (C) Correlation between titers against HerCOOH-BSA and SCR heroin ED\textsubscript{50} values collected over 26 weeks. P < 0.0001 by Pearson’s correlation. (D) 6AM serum concentrations over time in n = 4 rhesus macaques (M1–4) following 0.32 mg/kg i.m. heroin. The PK study was performed in the same subjects before and after a course of three initial immunizations. F\textsubscript{1,2,8} = 20.35, P = 0.0041 by a RM two-way ANOVA comparing pre and post vaccination PK (post-vacc 1). After 7 months, monkeys M1 and 2 received three additional immunizations and the PK study was repeated (post-vacc 2). (E) 6AM AUC values corresponding to the PK study. Fold increases versus baseline are reported as mean ± SEM. **P < 0.01 by paired t-test. C\textsubscript{max} values (ng/mL): baseline = 32.5 ± 6.8, post-vacc 1 = 418 ± 110, post-vacc 2 = 5430 ± 2800.
while titer is typically the metric used in gauging vaccine success, antibody affinity is also critical. The low nanomolar antibody affinity for 6AM, as observed by SPR, provides a competitive sink with brain MORs for drug binding based on Le Chatelier’s principle, thus causing heroin potency reduction in behavioral assays.

Heroin hapten design was greatly improved through conversion of the terminal thiol (HerSH) to a carboxylic acid (HerCOOH). This modification enabled more efficient and reliable protein coupling, producing a more efficacious immunoconjugate with higher epitope density. Furthermore, the HerCOOH hapten contains a shorter, more “immunologically silent” linker with less chemical functionalities relative to HerSH and HerdBA. Following antigen processing, the HerCOOH hapten linker likely interferes minimally with immune presentation of a heroin-like epitope. Previous studies have shown that peptidic linkers can increase antihapten immune responses,32−34 possibly by enhancing hapten anchoring to the MHCII (Figure 2). However, the dBA linker was not effective, which may be explained by the fact that beta-alanine is not naturally found in proteins, thus hampering immune processing and presentation of HerdBA. Because it was not clear whether heroin or 6AM should be emulated for linker was not e

10 In contrast, we found that cyclic dinucleotide adjuvants.67,68 These effects were synergistic with alum-mediated induction of humoral immunity, as shown previously.5,60,69,70 As only one dosage of CpG ODN and alum was investigated in the current study, fine-tuning of CpG ODN and alum dosing in future monkeys studies would be anticipated to further enhance vaccine efficacy.

Once a heroin vaccine formulation was identified that was efficacious in generating long-term, antibody-mediated protection from heroin PD, studies in rhesus macaques were pursued to evaluate the clinical potential of the vaccine. One major hurdle for drug vaccines in human trials is achieving efficacy in all patients. Although clinical trials for nicotine and cocaine vaccines have failed to demonstrate significant efficacy, they did demonstrate a proof-of-concept that patients who generated a strong antidrug antibody response displayed protection from the abuse-related drug effects.28−31 In order to improve drug vaccine performance, preclinical testing must employ clinically relevant benchmarks focusing directly on vaccine capacity to selectively alter the behavioral pharmacology of the target drug. Our studies in an outbred strain of mice (Swiss Webster) demonstrated that the vaccine reliably induced antidrug titers and significantly shifted the heroin dose–effect curves in the antinociception assay. In fact, the only detectable source of immune variability was the degree of hapten conjugation; heroin conjugates with greater haptenation produced greater efficacy upon immunization, which has been observed previously.56,71

The optimized vaccine formulation identified from the mouse studies translated well to rhesus monkeys, albeit with smaller heroin potency shifts, possibly due to immunological differences between species or vaccine dosages. Moreover, the large initial spike in vaccine response that was only observed in mice can be explained by the differing vaccine injection routes, i.p. in mice vs i.m. in monkeys, and in the former case, the vaccine can rapidly drain to the spleen to induce a strong, short-lived B-cell response. Regardless, significant antiheron IgG titers with low nanomolar 6AM affinity were observed in all four monkeys. In addition, the humoral immune response to the vaccine significantly altered heroin PK, increasing 6AM AUC values by ∼19-fold and half-life by ∼3-fold, resulting in the observed reduction in heroin PD. The PK results are in agreement with previous studies, which have noted that conjugate-vaccinated animals showed higher drug concentrations in serum due to antibody sequestration leading to lower drug concentrations in the brain relative to controls—a phenomenon observed for opioids65,66,72,73 and other drugs such as methamphetamine, cocaine and nicotine.74−77 In a manner that has not been fully studied, the drug-specific antibodies appear to increase drug half-life and diminish drug clearance. This is likely caused by antibody-drug binding, which hampers glomerular filtration of drugs by the kidneys and shields drugs from metabolic destruction. Eventual drug metabolism and clearance does not appear to be mediated by immunological mechanisms but rather by slow dissociation of antibody-drug complexes.72

... cells via TLR9 to produce an antidrug humoral response contributes to the success of CpG ODN 1826 and 2006 as adjuvants.67,68 These effects were synergistic with alum-mediated induction of humoral immunity, as shown previously.5,60,69,70 As only one dosage of CpG ODN and alum was investigated in the current study, fine-tuning of CpG ODN and alum dosing in future monkeys studies would be anticipated to further enhance vaccine efficacy.
three vaccinations spaced 6 weeks apart or greater is required for achieving significant immunity against heroin; however, as observed in monkeys M1,2, vaccination on an even longer time scale may be optimal. Importantly, no evidence of immune tolerance to the vaccine was seen at any point despite multiple immunization rounds and frequent administration of heroin as a result of SCR testing. The immunochemical and PK data support the heroin SCR data, the latter of which revealed an approximately 4-fold shift in the heroin dose–effect curve following each immunization. While the decrease in heroin potency was observed via the i.m. route of administration, these results are unlikely to be significantly different than the i.v. route commonly used by humans. The relative ratios of heroin, 6AM, and morphine after i.m. administration were consistent with the PK profile observed following i.v. heroin administration in humans.78 Furthermore, our first generation vaccine was effective in attenuating i.v. heroin self-administration in rats.77 Specifically, the vaccine could block heroin reinstatement following a single bolus of 0.18 mg/kg i.v. heroin, which would translate to a 14.4 mg dose in an 80 kg adult. This dose is within the dose range reported by heroin users (see Erowid.org). As our second generation vaccine has been shown to be at least 3-fold better than the first generation vaccine in the mouse antinociception model, we posit the second generation vaccine’s capacity to neutralize i.v. heroin would only be enhanced. Consolidation of our data strongly suggests that the vaccine acts as a heroin immunonantagonist; furthermore, the vaccine effects paralleled the effects of the FDA-approved opioid antagonist NTX. In previous rhesus monkey studies, NTX antagonized the rate suppressant, antinociceptive and discriminative stimulus effects of heroin.36,79,80 Similarly, reductions in heroin vs food choice have been demonstrated by administration of another opioid antagonist nalorex in nonopioid dependent monkeys81 and NTX pretreatment in nonopioid dependent rats.82 These results provide empirical evidence that the vaccine-mediated antagonism of heroin in the SCR assay would be predictive of antagonism of heroin effects in other models such as drug discrimination and self-administration. Given the noteworthy vaccine results in the SCR procedure, future studies to determine vaccine effects in more complex NHP behavioral procedures, e.g., i.v. self-administration, are warranted.

Other studies in NHPs have demonstrated the efficacy of optimized nicotine and cocaine vaccines. The second generation nicotine conjugate (NIC7-CRM) was optimized in mice83 and translated well to cynomolgus monkeys in a CpG ODN + alum formulation to elicit sustained (~10^6) antinocitine titers.84 Following a nicotine challenge in vaccinated monkeys, blood nicotine C_{max} and AUC increased by 29- and 89-fold, respectively, and nicotine brain concentrations were reduced.71 A fully synthetic nicotine vaccine (SEL-068) also translated from mice to monkeys83 and has been shown to attenuate nicotine discrimination in squirrel monkeys while reducing nicotine potency by 3-fold.86 The GNE cocaine hapten conjugated to a disrupted adenovirus (dAdS) and formulated with a proprietary adjuvant (Adjuplex) has consistently generated high anticocaine titers (10^6–10^8) in a number of rhesus macaque studies.87–90 This vaccine increased cocaine C_{max} in serum by ~3.5-fold, mitigated biodistribution of cocaine to the brain and other organs,87,88,89 and attenuated cocaine self-administration and reacquisition.90 Although these reports provide some context for our NHP heroin vaccine studies, direct comparisons between our studies and others are limited. In all cases, antihapten titer levels responded similarly to the vaccines; however, our studies use midpoint titers, which are intrinsically 10-100 fold lower than the end point titers used in other studies. Furthermore, hapten-specific titer levels do not necessarily reflect affinity to the actual target drug, and thus are not always indicative of efficacy, although a robust correlation between titer and efficacy was observed for the disclosed heroin vaccine. While previous NHP vaccine studies have not fully investigated drug affinity of monkey antiserum, we have observed 0.5–6 nM affinity to 6AM. Comparatively, dAdSNGNE in NHPs was shown to produce 5–120 nM antibody affinity to cocaine.87 In considering behavioral testing, few studies have compared drug ED_{50} in vaccinated and nonvaccinated animals whereas we have demonstrated heroin dose–effect curve shifting in both mouse and monkey models. For reference, the SEL-068 vaccine reduced nicotine potency by 3-fold in monkeys while we saw a 4-fold reduction in heroin potency. However, results in the more complex self-administration procedures used in a cocaine vaccine study94 and the first heroin vaccine study,24 cannot be related to the ED_{50} determinations in our study. On the other hand, PK metrics in vaccinated monkeys can be compared with the caveat that drug PK/PD between each of the drugs differ drastically. Vaccine-mediated fold-increases in heroin C_{max} for HerCOOH-TT (post-vacc 1) appeared to be half that of the nicotine C_{max} for NIC7-CRM34 but 4-fold greater than cocaine C_{max} for dAdSNGNE95 therefore, our vaccine possesses excellent serum neutralizing capacity for heroin and its psychoactive metabolites relative to other vaccines.

In response to the opioid epidemic that has plagued the United States, the disclosed heroin vaccine could satisfy a dire, unmet need for an opioid use disorder therapeutic. Since the vaccine reduces heroin potency, vaccinated drug users would encounter an increased cost to “getting high”, potentially similar to the effects of clinically available and FDA-approved depot NTX formulations.91,92 Indeed, our previous studies have shown that following a period of drug abstinence, drug-dependent, immunized rats actually extinguish self-administration of heroin,37 and these rats demonstrated 4-fold shifted heroin dose–effect curves in antinociceptive testing. Considering this study and the similar curve shifts observed in monkeys, our vaccine has the potential to be clinically useful; it may serve as a “safety catch” for preventing relapse episodes in former heroin users attempting to maintain drug abstinence, or it may help current heroin abusers to achieve abstinence. This hypothesis is supported by clinical studies showing that long-acting depot NTX mitigated the reinforcing effects of heroin in humans,25,93 and instead of heroin users increasing their drug intake to surmount the antagonist, heroin extinction-like behaviors occurred.91,94 The obvious major benefits to vaccination over pharmacological antagonists are the potential for increased duration of action and decreased side effects. Coadministration of opioid agonists methadone or buprenorphine as needed along with the vaccine would likely enhance therapeutic efficacy by alleviating opioid cravings. Such combination therapy would be possible due to the vaccine’s selective sequestration of 6AM by >1000-fold over other opioids; moreover, prescription pain medication, e.g., oxycodone also would not interact with the heroin vaccine.

In conclusion, an efficacious heroin vaccine has been identified through optimization of the adjuvant (CpG ODN + alum), carrier protein (TT) and hapten (HerCOOH). The
vaccine is efficacious in basic preclinical mouse and NHP models over a wide range of heroin doses, accomplishing an important milestone in the drug development process to human clinical trials. Forty years after the first report of a heroin/morphine vaccine, numerous studies and research groups have advanced the concept of antidrug vaccines to a functional level, whereby the vaccines act as a long-term “immunoantagonist” to attenuate drug PD. While we have speculated as to how our particular heroin vaccine could be used to treat use disorder, future studies involving more advanced NHP models and clinical trials must be performed to elucidate the true therapeutic utility of this vaccine as well as other opioid vaccines.

**ASSOCIATED CONTENT**

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b03334.

Materials and methods, conjugate mass data, titer data, additional mouse and monkey vaccine data: Tables S1–S2, Figures S1–S10 (PDF)

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