Novel Molecule Exhibiting Selective Affinity for GABA<sub>A</sub> Receptor Subtypes

Cecilia M. Borghese<sup>1</sup>, Melissa Herman<sup>2,6</sup>, Lawrence D. Snell<sup>3</sup>, Keri J. Lawrence<sup>4</sup>, Hyun-Young Lee<sup>1</sup>, Donald S. Backos<sup>6</sup>, Lauren A. Vanderlinden<sup>4</sup>, R. Adron Harris<sup>1</sup>, Marisa Roberto<sup>7</sup>, Paula L. Hoffman<sup>3,5</sup> & Boris Tabakoff<sup>3,6</sup>

Aminoquinoline derivatives were evaluated against a panel of receptors/channels/transporters in radioligand binding experiments. One of these derivatives (DCUK-OEt) displayed micromolar affinity for brain GABA<sub>γ</sub>-aminobutyric acid type A (GABA<sub>A</sub>) receptors. DCKU-OEt was shown to be a positive allosteric modulator (PAM) of GABA currents with α1β2-γ2, α1β3-γ2, α5β3-γ2 and α1δ3-γ6 GABA<sub>A</sub> receptors, while having no significant PAM effect on α1β3 receptors or α1β2-γ1, α3β3-γ2 or α4β3-γ6 receptors. DCKU-OEt modulation of α1β2-γ2 GABA<sub>A</sub> receptors was not blocked by flumazenil. The subunit requirements for DCKU-OEt actions distinguished DCKU-OEt from other currently known modulators of GABA function (e.g., anesthetics, neurosteroids or ethanol). Simulated docking of DCKU-OEt at the GABA<sub>A</sub> receptor suggested that its binding site may be at the α + β subunit interface. In slices of the central amygdala, DCKU-OEt acted primarily on extrasynaptic GABA<sub>A</sub> receptors containing the α1 subunit and generated increases in extrasynaptic “tonic” current with no significant effect on phasic responses to GABA. DCKU-OEt is a novel chemical structure acting as a PAM at particular GABA<sub>A</sub> receptors. Given that neurons in the central amygdala responding to DCKU-OEt were recently identified as relevant for alcohol dependence, DCKU-OEt should be further evaluated for the treatment of alcoholism.

GABA (γ-aminobutyric acid) is the major inhibitory transmitter and glutamate is the major excitatory transmitter in brain and these two opposing forces are in constant interplay within the communication systems of the brain<sup>1</sup>. The desire for pharmacological manipulation of GABAergic neurotransmission has generated a plethora of xenobiotics which are useful in medicine, including anticonvulsants, anesthetics, anxietytics, muscle relaxants and medications for treating pain. The realization that the GABA<sub>A</sub> receptor system is a collage derived from 6 α, 3 β, 3 γ<sub>1</sub>, 6, 6, 6, 6, 6 and 3 θ subunits<sup>2,3</sup>, and that different combinations of these subunits are particularly important in certain physiologic events mediated by GABA<sub>A</sub>, has stimulated a search for chemical entities that have selectivity for GABA<sub>A</sub> receptors with a particular combination of subunits.<sup>2,5</sup>

We had previously reported on a “rationally engineered” molecule which effectively reduced allodynia in animal models of neuropathy by simultaneously targeting the NMDA subtype of glutamate receptor and voltage-sensitive sodium channels<sup>4</sup>, particularly Na<sub>1.7</sub><sup>4</sup> and Na<sub>1.8</sub><sup>8</sup>. This compound showed neither sedative effects per se, nor did it enhance the sedative or motor incoordinating effects of ethanol. We more recently generated a number of chemical derivatives of the “skeleton quinoline structure” of our original molecule. In screening these molecules through a series of radioligand binding assays<sup>9</sup> we found that 5,7-dichloro-4-[(diphenyl carbamoyl) amino] quinoline-2-ethyl carboxylate (DCUK-OEt) (Fig. 1) could displace muscimol from its specific binding sites in an assay containing washed rat brain membranes, while it had no effect at a concentration of 10 μM in 32 other radioligand binding assays. The current manuscript describes the equilibrium radioligand binding studies and electrophysiological analysis of the effects of DCKU-OEt, as well as the non-esterified derivative, 5,7.

<sup>1</sup>The University of Texas at Austin, Waggoner Center for Alcohol and Addiction Research, Austin, TX, 78712, USA. 2Department of Neuroscience, The Scripps Research Institute, La Jolla, CA, 92037, USA. 3Lohola Research Corporation, Aurora, CO, 80045, USA. 4Department of Pharmaceutical Sciences, University of Colorado Skaggs School of Pharmacy & Pharmaceutical Sciences, Aurora, CO, 80045, USA. 5Department of Pharmacology, University of Colorado School of Medicine, Aurora, CO, 80045, USA. 6Present address: Bowles Alcohol Research Center, Chapel Hill, NC, 27514, USA. Correspondence and requests for materials should be addressed to B.T. (email: boris.tabakoff@ucdenver.edu)
Figure 1. Chemical structure of DCUK compounds. (a) DCUKA (5,7-Dichloro-4-((diphenyl carbamoyl) amino) quinoline-2-carboxylic acid). (b) DCUK-OEt (5,7-Dichloro-4-((diphenyl carbamoyl) amino) quinoline-2-ethyl carboxylate).

dichloro-4-((diphenyl carbamoyl) amino) quinoline-2-carboxylic acid (DCUKA) (Fig. 1) which is the primary metabolite of DCUK-OEt, on GABA\textsubscript{A} receptors. The electrophysiological studies were carried out in *Xenopus laevis* oocytes and in neurons from the rat central amygdala (CeA). The GABA\textsubscript{A} subunit combinations tested in oocytes were selected based on their abundance in brain (e.g. α1/β2-2) and their expression in the CeA\textsuperscript{10-14}. Additional subunits were expressed with the objective of further elucidating the selectivity of the DCUK compounds. The results indicate that DCUK-OEt may have characteristics which distinguish it from all currently available ligands that act on the GABA\textsubscript{A} receptor.

**Results**

The radioligand displacement studies that were performed with \(^{3}H\)flunitrazepam and \(^{3}H\)muscimol, utilized washed rat brain membranes and thus represented an amalgam of GABA\textsubscript{A} receptors composed of various subunit combinations. Neither DCUK-OEt nor DCUKA demonstrated efficacy for displacing \(^{3}H\)flunitrazepam. However, at concentrations <10 \(\mu\)M, both DCUK-OEt and DCUKA were able to displace \(^{3}H\)muscimol, albeit with different potency. The \(K_{i}\) for displacement of muscimol binding by DCUKA was 6.6 \(\mu\)M and displacement by DCUK-OEt demonstrated a lower \(K_{i}\) of 1.7 \(\mu\)M (Table 1). DCUK-OEt at concentrations <10 \(\mu\)M demonstrated no significant displacement of any of the ligands selective for 32 other receptors/transporters/channels that were tested in the course of our studies (Supplementary Table S1).

Both DCUK-OEt and DCUKA enhanced submaximal GABA (EC\textsubscript{10}) currents in oocytes containing α1/β2-2 GABA\textsubscript{A} receptors (Fig. 2a). Full concentration-response curves were not possible due to solubility limits, but, from the partial curves, equi-effective concentrations were approximately 10-fold lower for DCUK-OEt than for DCUKA (e.g., 0.3 \(\mu\)M DCUK-OEt had the same effect as 3 \(\mu\)M DCUKA). DCUK-OEt was similarly effective in potentiating submaximal GABA currents in α1/β3 and α1/β2-2 GABA\textsubscript{A} receptors (Fig. 2a and b). Interestingly, DCUK-OEt potentiated GABA currents produced by higher concentrations of GABA (EC\textsubscript{60} and EC\textsubscript{100}) with α1/β3 GABA\textsubscript{A} receptors, but not with α1/β2-2 GABA\textsubscript{A} receptors (Fig. 2c). Representative tracings of GABA-induced currents in the presence of DCUK-OEt are shown in Supplementary Fig. S1. The positive modulation of GABA\textsubscript{A} receptors by DCUK-OEt was specific to the GABA\textsubscript{A} family of heteromeric receptors and even closely related receptors such as \(\rho\) GABA\textsubscript{A} and \(\alpha\) Gly receptors showed no evidence of positive allosteric modulator (PAM) activity with DCUK-OEt (DCUK-OEt produced a small but statistically significant reduction in \(\rho\)1 receptor currents, Supplementary Fig. S2).

A prominent group of positive allosteric modulators of GABA\textsubscript{A} receptors act through the benzodiazepine site, located at the extracellular interface between the α and the γ subunit\textsuperscript{4}. To test whether the DCUK compounds act through this site, we co-applied flumazenil. Flumazenil can act as a partial agonist at the benzodiazepine site, and 20 \(\mu\)M flumazenil alone potentiated EC\textsubscript{10} GABA responses (51 ± 2\%, \(n = 5\)). However, flumazenil did not significantly affect either DCUK-OEt (Fig. 2d) or DCUKA (not shown) actions on α1/β2-2 GABA\textsubscript{A} receptors, while significantly inhibiting flunitrazepam PAM actions. In these studies, 0.1 \(\mu\)M flumazenil showed no significant displacement of any of the ligands selective for 32 other receptors/transporters/channels that were tested in the course of our studies (Supplementary Table S1).

The composition of the GABA\textsubscript{A} receptors was critical in determining the effects of DCUK-OEt. When applied to α1/β2, α5/δ3 or α1/δ3 GABA\textsubscript{A} receptors, the average effect of 0.3 \(\mu\)M DCUK-OEt was not significantly different from zero (Table 2). A third subunit (either γ or δ) definitively increased the DCUK-OEt PAM effect, and the identity of the third subunit was quite relevant to the magnitude of the PAM effect. For instance, DCUK-OEt induced less potentiation (non-significant) of the GABA responses with α1/β2-3 GABA\textsubscript{A} receptors compared with α1/β2-2 GABA\textsubscript{A} receptors (Tables 2 and 3).

The identity of the α subunit also contributed to the magnitude of the DCUK-OEt effect: DCUK-OEt significantly potentiated GABA responses of α1/β3, but not α4/β3 GABA\textsubscript{A} receptors (Tables 2 and 3). DCUK-OEt similarly potentiated α1/β3-2 and α5/δ3-2 GABA\textsubscript{A} receptors, but the PAM effect was not significantly different from zero for α4/δ3-2 GABA\textsubscript{A} receptors (Table 2). The identity of the β subunit also played a role in the magnitude
of the DCUK-OEt effect as a PAM: the $\alpha_1\beta_1\gamma_2$ GABA$_A$ receptors showed no significant potentiation of the GABA responses by DCUK-OEt while $\alpha_1\beta_2\gamma_2$ and $\alpha_1\beta_3\gamma_2$ GABA$_A$ receptors did (Table 2). The $\beta_1$ subunit residue 265 seems to play an important role in determining the effect of certain GABA$_A$ receptor modulators: when S265 in $\beta_1$ is mutated to N (homologous residue in $\beta_2$ and $\beta_3$) on the GABA$_A$ receptor complex, the modulators’ potentiation is increased, and vice versa, when N265 in $\beta_2$ or $\beta_3$ is mutated to S, the potentiation is reduced$^{16–18}$. When we tested DCUK-OEt on $\alpha_1\beta_2(N265S)\gamma_2$ compared to $\alpha_1\beta_2\gamma_2$, the effect of DCUK-OEt as a PAM was significantly reduced (Table 3), but not to the extent seen with drugs such as etomidate (no GABA potentiating effect of etomidate at concentrations up to 1 mM was evident with the $\alpha_1\beta_2(N265S)\gamma_2$ receptor combination) $^{16}$.

To further investigate potential binding sites for DCUK-OEt on the GABA$_A$ receptor, we performed computationally-based small molecule docking studies to compare the potential interactions of DCUK-OEt with those of DCUKA, flunitrazepam, and etomidate, with either the classical benzodiazepine binding site (located at the $\alpha + \beta$ interface of the pentameric receptor) or an alternative binding site (at the $\alpha + \beta$ interface) (Fig. 3a).

The corresponding binding energies are shown in Table 4. These studies indicated that DCUK-OEt exhibited the highest predicted affinity for an alternative binding site, while, as expected, flunitrazepam exhibited the highest predicted affinity for the benzodiazepine site.

The modeling studies predicted both the carboxylate of DCUKA and the ethyl ester moiety of DCUK-OEt to be oriented towards the $\alpha$ subunit in the region of $\alpha$-Tyr160 in the alternative site (Fig. 3b and c). The ethyl ester was predicted to participate in additional hydrophobic interactions with the residues of this region, and there exists a potential $\pi$-$\sigma$ interaction with $\alpha$-Tyr160. These additional interactions of the ethyl ester also appeared to

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**Table 1.** Displacement of Ligands Binding to GABA$_A$ Receptors by DCUK-OEt and DCUKA. IC$_{50}$ and Ki values were obtained by non-linear regression analysis of radioligand binding isotherms. Ki values are reported as estimates from the non-linear regressions and their associated standard errors (n = 10 points in the binding isotherms).

| Compound | [3H]Muscimol Binding | [3H]Flunitrazepam Binding |
|----------|----------------|--------------------------|
| DCUK-OEt | 1.7 ± 0.3 µM | >10 µM |
| DCUKA | 6.6 ± 1.9 µM | >10 µM |

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Figure 2. DCUK effect on GABA responses. (a) Effect of DCUK compounds on submaximal (EC$_{10}$) GABA responses of $\alpha_1\beta_2\gamma_2$ GABA$_A$ receptors (n = 4–5 at each concentration of DCUK compound). (b) Effect of DCUK-OEt on submaximal (EC$_{10}$) GABA responses of $\alpha_1\beta_3\gamma_2$ GABA$_A$ receptors (n = 5–6 at each concentration of DCUK-OEt). (c) Effect of DCUK-OEt (0.3 µM) and escalating GABA concentrations applied to $\alpha_1\beta_2\gamma_2$ and $\alpha_1\beta_3\gamma_2$ GABA$_A$ receptors (n = 9 each). GABA concentrations used: 3 and 1 µM for $\alpha_1\beta_2\gamma_2$ and $\alpha_1\beta_3\gamma_2$, respectively (~EC$_{10}$); 30 µM (~EC$_{60}$); 3 mM (~EC$_{100}$). (d) DCUK-OEt (0.3 µM) effect in the absence and presence of 20 µM flumazenil (Flu) (n = 5 for each condition). Data represent mean ± SEM. **p < 0.01 compared to $\alpha_1\beta_2\gamma_2$ (one-way ANOVA and post-hoc contrasts).
optimize the positioning of the head group and amide linker within the binding pocket to allow for additional potential H-bond and π-π interactions with β:Asp43 and Tyr62, respectively, leading to the higher affinity for DCUK-OEt compared to DCUKA for the GABA receptor.

The predicted binding and interactions of flunitrazepam in the benzodiazepine site (Supplementary Fig. S4) were consistent with previous studies, and flunitrazepam made a number of favorable contacts, including H-bond interactions with α:Tyr160 and γ:Tyr142, and π-π interactions with α:Tyr160 and Tyr210. DCUKA shared a number of these predicted contacts, while the ethyl ester of DCUK-OEt appeared to impair the optimal positioning of the head group in the benzodiazepine binding pocket (Fig. 3d and e). Flunitrazepam bound somewhat more deeply into the pocket, compared to the other tested compounds, with the fluorobenzene ring predicted to be locked in place by a three-way π-stacking interaction with α:His102 and γ:Tyr58, Phe77. An additional π-σ interaction with α:Phe100 and π-π stacking with γ:Phe77 not only distinguished the predicted binding of flunitrazepam from DCUKA, but also represent the crucial aspects of interaction of flunitrazepam with the receptor.

### Table 2. DCUK-OEt (0.3 μM) induced change in the response to EC₅₀ GABA in GABA₃ receptors composed of different subunit combinations. Significant and marginal effects are those with a Bonferroni-adjusted p-value < 0.05 and < 0.2, respectively. A linear mixed model was implemented in SAS (version 9.4) to calculate the normalized percent change in current for each receptor subunit combination produced by DCUK-OEt (EC₅₀ GABA concentration without and with 0.3 μM DCUK-OEt). A random effect of batch was included in the model, and for each receptor, the percent change in the GABA-induced current produced by DCUK-OEt was compared to 0 using a single-sample t-test in the MIXED procedure in SAS and a Bonferroni adjustment to correct for multiple comparisons.

| Receptor | Percent Change | Standard Error | Sample Size | Unadjusted p-value | Bonferroni adjusted p-value | Effect |
|----------|----------------|----------------|-------------|--------------------|----------------------------|--------|
| α1/2     | 40             | 19             | 11          | 0.035              | 0.42                       | Non-significant |
| α1/3     | 20             | 27             | 5           | 0.449              | >0.99                      | Non-significant |
| α3/3     | 8              | 25             | 8           | 0.730              | >0.99                      | Non-significant |
| α1/1+2   | 17             | 23             | 11          | 0.445              | >0.99                      | Non-significant |
| α1/2+2   | 127            | 10             | 44          | <0.001             | <0.01                      | Significant   |
| α1/2(N265S)+2 | 45       | 18             | 14          | 0.013              | 0.16                       | Marginal     |
| α1/2+1   | 56             | 24             | 8           | 0.022              | 0.26                       | Non-significant |
| α1/3+2   | 95             | 29             | 6           | 0.002              | 0.02                       | Significant   |
| α4/3+2   | 49             | 26             | 8           | 0.066              | 0.79                       | Non-significant |
| α5/3+2   | 81             | 24             | 8           | 0.001              | 0.01                       | Significant   |
| α1/38    | 102            | 16             | 18          | <0.001             | <0.01                      | Significant   |
| α4/38    | -6             | 31             | 5           | 0.842              | >0.99                      | Non-significant |

### Table 3. Comparison of DCUK-OEt induced changes in EC₅₀ GABA responses between receptors differing in a single subunit. These comparisons were executed with the linear mixed model using linear contrasts. Correction for multiple pairwise comparisons was by a Bonferroni adjustment.

| Receptor 1 | Receptor 2 | Percent Difference (Receptor 1-2) | Standard Error | Unadjusted p-value | Bonferroni adjusted p-value | Effect |
|------------|------------|----------------------------------|----------------|--------------------|----------------------------|--------|
| α1/2       | α1/3       | -87                              | 19.1           | <0.01              | <0.01                      | Significant |
| α1/2+2     | α1/3+2     | 110                              | 24.7           | <0.01              | <0.01                      | Significant |
| α1/2+2     | α1/3+2     | 32                               | 30.9           | 0.30               | >0.99                      | Non-significant |
| α1/2+2     | α1/3(N265S)+2 | 82   | 17.5                          | <0.01          | <0.01                      | Significant |
| α1/2+2     | α1/3(N265S)+2 | -28  | 28.7                          | 0.34            | >0.99                      | Non-significant |
| α1/2+2     | α1/3+2     | -77                              | 36.9           | 0.04               | 0.69                       | Non-significant |
| α1/2+2     | α1/3+2     | 71                               | 24.6           | <0.01              | 0.09                       | Marginal |
| α1/2       | α1/3+3     | 20                               | 32.5           | 0.54               | >0.99                      | Non-significant |
| α1/3+3     | α5/3       | 12                               | 36.3           | 0.74               | >0.99                      | Non-significant |
| α1/3       | α1/3+3     | -74                              | 39.6           | 0.06               | >0.99                      | Non-significant |
| α5/3       | α5/3+2     | -72                              | 34.5           | 0.04               | 0.69                       | Non-significant |
| α1/3+2     | α4/3+3     | 46                               | 59.2           | 0.24               | >0.99                      | Non-significant |
| α1/3+2     | α3/3+2     | 14                               | 38.0           | 0.71               | >0.99                      | Non-significant |
| α4/3+2     | α3/3+2     | -32                              | 35.7           | 0.37               | >0.99                      | Non-significant |
| α1/3       | α1/38      | -82                              | 27.9           | <0.01              | 0.07                       | Marginal |
| α1/3+2     | α1/38      | -8                               | 33.2           | 0.82               | >0.99                      | Non-significant |
| α4/3+2     | α4/38      | 55                               | 36.8           | 0.14               | >0.99                      | Non-significant |
| α1/3+2     | α4/38      | 109                              | 35.0           | <0.01              | 0.05                       | Significant |
that lead to its pharmacological function. The presence of the phenyl (C ring) substitution at the 5 position of the benzodiazepine ring structure is necessary for the PAM actions of the benzodiazepine derivatives 22, 23. Therefore, even though DCUKA and DCUK-OEt may bind to the benzodiazepine binding site on the GABA_A receptor (with lower affinity), the lack of the fluorbenzene ring on the DCUKA and DCUK-OEt structures would predict the lack of functional effect of DCUKA and DCUK-OEt via the benzodiazepine site. It is important to note that, due to the method by which binding energies are calculated, comparisons of relative binding affinity can only be reliably assessed between different molecules within the same binding site.

The studies showing that potentiation of GABA responses by DCUK-OEt cannot be blocked by flumazenil do not preclude the possibility, suggested by the docking experiments, that DCUK-OEt could bind to the benzodiazepine site as an antagonist, while producing potentiation via binding to a different site (the alternative, extracellular site or a transmembrane one). We tested this hypothesis by co-applying DCUK-OEt (1 µM) and

Figure 3. Predicted docking of DCUK-OEt and DCUKA within extracellular domain interfaces of GABA_A receptor subunits. The α subunit is shaded in green, β in cyan and γ in yellow. (a) Extracellular (top down) view of the pentameric GABA_A receptor. The interfaces illustrated are α + β- (alternative site), and α + γ- (benzodiazepine site). (b) DCUK-OEt and (c) DCUKA within the alternative site (α + β-). DCUK-OEt is represented by orange sticks and DCUKA is represented by pink sticks. (d) DCUKA and (e) DCUK-OEt within the benzodiazepine site (α + γ-). Dashed lines indicate predicted non-bond interactions (green = H-bonds, orange = electrostatic or π-cation/anion, magenta = π-π, purple = π-σ, pink = hydrophobic).
Flunitrazepam (0.1 μM). The combined effect was larger than the sum of their individual effects (Supplementary Fig. S5), suggesting that the functional effects of the two drugs may be mediated by actions at two different sites.

The significant effects of DCUK-OEt on particular subunit combinations of the GABA_A receptor led us to test the effects of this compound on neurons in the rat central amygdala (CeA). The CeA is primarily composed of GABAergic neurons and changes in CeA GABAergic neurotransmission have been implicated in the development and maintenance of alcohol dependence. Focal application of DCUK-OEt (0.5 μM) was found to reduce the holding current in medial CeA neurons (Fig. 4a and b), while producing no significant effect on spontaneous inhibitory postsynaptic current (sIPSC) frequency, amplitude, rise or decay times (Fig. 4c).

To confirm that the changes in holding current that we observed were due to increases in tonic signaling at the GABA_A receptor, the GABA_A receptor antagonist gabazine (GBZ, 100 μM, Sigma Chemical Co., St. Louis, MO) was focally applied following DCUK-OEt application. GBZ produced a significant reduction in holding current when applied after DCUK-OEt, suggesting that the changes in holding current that were observed with DCUK-OEt were due to DCUK-OEt-induced increases in tonic conductance via GABA_A receptors on medial CeA neurons. In addition, we found that the increase in holding current with DCUK-OEt was positively correlated with the reduction in holding current seen with GBZ application (Pearson correlation coefficient = 0.838; p = 0.0094; n = 8; Fig. 4d).

**Discussion**

DCUK-OEt acts as a subunit-selective PAM at the GABA_A receptor, and our ligand binding studies produced no evidence of interaction of DCUK-OEt (<10 μM) with 32 other receptors/transporters/channel proteins. DCUK-OEt exhibited its most robust effects on submaximal GABA-induced currents when applied to the α1β2-δ GABA_A receptor, the subunit combination most highly expressed in mammalian brain. Similar PAM activity of DCUK-OEt was observed with GABA_A receptors composed of α1β3δ subunits. On the other hand, DCUKA, which lacks the ester moiety at the 2 position of the carbonylquinoline, and is the major metabolite of DCUK-OEt, was 10 times less potent than DCUK-OEt in acting as a PAM at the GABA_A receptor.

The most studied PAMs at the GABA_A receptor are benzodiazepine derivatives and other compounds (e.g., zolpidem) which act at the interface of extracellular domains of the α and γ subunits. Our data produced no evidence for DCUK-OEt action at this site. DCUK-OEt did not displace flunitrazepam in ligand displacement experiments, and the electrophysiological effects of DCUK-OEt (and DCUKA) on GABA_A receptors expressed in oocytes were not modified by the selective benzodiazepine antagonist, flumazenil. Additionally, the substitution of a δ subunit for a γ subunit in the GABA_A receptor complex greatly diminishes the effects of benzodiazepines but the effects of DCUK-OEt were similar in receptors containing either γ2 or δ subunits (compare α1β3γ2 and α1β3δ in Tables 2 and 3). Finally, when DCUK-OEt and flunitrazepam were applied together, the PAM effect was supra-additive.

Table 4. Docking binding energies and interactions at GABA_A receptor sites. Summary of the binding energies and non-bond interactions of the top scoring predicted binding orientations for each compound docked into the homology model of the benzodiazepine binding site at the α+γ subunit interface or the “Alternative” binding site at the α+β subunit interface of the human GABA_A receptor shown in Fig. 2 and in Supplementary Fig. S4. Binding orientations were predicted using the Discovery Studio flexible docking protocol and energies were calculated using the distance-dependent dielectric model, as outlined in the methods.
site\textsuperscript{26} are particularly effective as PAMs, and are also direct agonists at GABA\textsubscript{A} receptors composed of the $\alpha_4$/$\beta_6$ subunits, while DCUK-OEt had no significant effect on this subunit combination. Furthermore, the modulatory action of neurosteroids at low concentrations does not differ among $\beta$ subunits\textsuperscript{27}. On the other hand, both $\beta_2$ and $\beta_3$ subunits in combination with $\alpha_1$ and $\gamma_2$ subunits responded to the addition of DCUK-OEt with a significant increase in the current induced by submaximal GABA, but the substitution of the $\beta_1$ subunit for either $\beta_2$ or $\beta_3$ resulted in a notable decrease of the PAM activity of DCUK-OEt (Table 2). The negative effect of the $\beta_1$ subunit is reminiscent of the selectivity for $\beta$ subunits shown by modulators such as loreclezole \textsuperscript{18} and etomidate\textsuperscript{28, 29}, among others. Three amino acids in the transmembrane domains of the $\beta$ subunit, distinguish the sequence of $\beta_1$ from $\beta_2$/$\beta_3$\textsuperscript{30}, and mutation of the asparagine at position 265 in the $\beta_2$ sequence, located at the interface of $\alpha$/$\beta$ transmembrane domains, has been demonstrated to interfere with the potentiating action of etomidate and other anesthetics at GABA\textsubscript{A} receptors\textsuperscript{16, 17, 30, 31}. The introduction of a mutated $\beta_2$ (N265S) into a complex containing $\alpha_1$ and $\gamma_2$ subunits significantly reduced (Table 3) the PAM activity of DCUK-OEt. However, this mutation has been shown to eliminate etomidate's PAM action\textsuperscript{28, 32}. Mutation of $\beta_2$N265 also decreases alcohol PAM activity on GABA\textsubscript{A} receptors\textsuperscript{33, 34}. However, ethanol potentiates GABA effects at receptors composed of dimeric $\alpha$/$\beta$ GABA\textsubscript{A} receptors, and does not discriminate between $\beta_1$ versus $\beta_2$ subunits\textsuperscript{35}. Reports on the concentrations of ethanol necessary to potentiate the effects of GABA on $\alpha_4$/$\beta_6$ GABA\textsubscript{A} receptors expressed in Xenopus oocytes have been contradictory\textsuperscript{36–38}, but the ethanol effect on the $\alpha_4$/$\beta_6$ subunit combination is always potentiation of the GABA actions, in contrast to the lack of any significant effect of DCUK-OEt.

Figure 4. DCUK-OEt potentiates tonic currents in medial CeA neurons. (a and b) Focal application of DCUK-OEt (0.5 $\mu$M) significantly increased the holding current in medial CeA neurons (*p < 0.05, paired t-test). (c) No change was evident in frequency, amplitude, rise and decay of mIPSPs with focal application of DCUK-OEt. (d) Correlation of magnitude of increase in tonic current produced by 0.5 $\mu$M DCUK-OEt with reduction of current by subsequent application of 100 $\mu$M gabazine. To demonstrate that changes in holding current were due to increases in tonic signaling, the GABA\textsubscript{A} receptor antagonist gabazine (GBZ) (100 $\mu$M) was focally applied following DCUK-OEt application. For all graphs, n = 11 cells.
At the EC\textsubscript{10} concentration of GABA, DCUK-OEt exhibited PAM effects on \(\alpha\)1\beta\gamma\delta\ GABA\textsubscript{A} receptors similar to effects seen with \(\alpha\)1\beta2. However, DCUK-OEt also enhanced the current produced by saturating concentrations of GABA within the \(\alpha\)1\beta2\gamma\delta subunit combination, but not with the \(\alpha\)1\beta2\gamma2\delta2 combination (Fig. 2c). GABA has been shown to be a partial agonist at \(\delta\) subunit-containing receptors\textsuperscript{39,40}, and DCUK-OEt, and some other PAMs\textsuperscript{40}, may allow for further activation of the GABA\textsubscript{A} receptor at concentrations seemingly maximal in the absence of PAMs. It also should be stressed that we detected no effect of DCUK-OEt at any concentration on any of the subunit combinations we tested in our paradigm, without the addition of GABA.

Overall, as noted above, there seems to be some overlap in the characteristics of DCUK-OEt with properties exhibited by allopregnanolone, CGS 9895, LAU-177\textsuperscript{41,42}, loreclezole, etomidate and ethanol, but other characteristics regarding subunit selectivity of DCUK-OEt mitigate against assuming that DCUK-OEt binding/activity occurs specifically through the currently described site(s) for binding of these agents. Additionally, DCUK-OEt characteristics do not conform to what would be expected if DCUK-OEt were utilizing the canonical barbiturate, or intravenous or inhalation anesthetic sites to affect GABA action at the GABA\textsubscript{A} receptor\textsuperscript{36,43-45}.

Our models to ascertain the docking of DCUK-OEt to interfaces between the various subunits of the GABA\textsubscript{A} receptor (composed of \(\alpha\)1\beta2\γ2 subunits), indicated that a binding site for DCUK-OEt may exist between the \(\alpha + \beta\) interface in the pentameric receptor. The free energy (\(\Delta G\)) of binding at this site was highest for DCUK-OEt and lowest for etomidate and flunitrazepam. When examining the docking at the benzodiazepine site located between the \(\alpha + \beta\) interface, the order was reversed, with flunitrazepam showing the highest binding energy and DCUK-OEt and etomidate showing the lowest \(\Delta G\). If the function of DCUK-OEt was dependent on binding at a single site at the \(\alpha + \beta\) interface, one would expect that GABA\textsubscript{A} receptors composed of only \(\alpha\) and \(\beta\) subunits would respond as well as the receptors which also contain the \(\gamma\) or \(\delta\) subunit. This was not the case, and the presence of the \(\gamma\) or \(\delta\) subunit was necessary to exhibit the PAM action of DCUK-OEt. In fact, the type of \(\gamma\) subunit expressed with the \(\alpha\) and \(\beta\) subunits was important, with the \(\gamma1\) subunit being significantly less effective than the \(\gamma2\) subunit. Because of the absence of the phenyl ring substituent (C ring) that generates functional (PAM) benzodiazepine derivatives, DCUK-OEt would not be expected to be an agonist at the benzodiazepine site, and our electrophysiologic experiments in the presence of flumazenil support this contention. It was, however, interesting that the combined effects of flunitrazepam and DCUK-OEt produced significantly more than an additive effect, possibly indicating an allosteric interaction between the benzodiazepine site and the site on the \(\alpha + \beta\) interface which binds DCUK-OEt with higher affinity.

The radioligand binding studies that led us to the electrophysiological examination of DCUK-OEt on the GABA\textsubscript{A} receptor, also produced some insight into the possible mechanism by which DCUK-OEt may generate its effects. DCUK-OEt produced a decrease in the affinity for muscimol at the GABA\textsubscript{A} receptor. Such action may be expected if DCUK-OEt is shifting the GABA\textsubscript{A} receptor into a state more likely to be in an open channel configuration. The GABA\textsubscript{A} receptor has been shown to display two affinity states for agonists such as muscimol\textsuperscript{39,47} and the high affinity state of the GABA\textsubscript{A} receptor has been proposed to represent stabilization of the desensitized form of the receptor\textsuperscript{41,42}. One can speculate that DCUK-OEt is increasing the proportion of receptors in a low affinity state at any particular concentration of agonist (muscimol). This speculation will require more investigation, but it is interesting that ethanol\textsuperscript{32} and the anxiolytic/anticonvulsant etifoxine\textsuperscript{33}, both which can act as PAMs at lower concentrations, reduce muscimol affinity at GABA\textsubscript{A} receptor in rat brain membrane preparations.

The \(\alpha\)1\beta2\gamma\delta combination is the primary combination of synaptically localized GABA\textsubscript{A} receptors in brain that mediate phasic inhibition, while \(\alpha\)1\beta4\gamma\delta\ or \(\alpha\)1\beta4\gamma\delta\ receptors have been considered to be the primary type of extrasynaptic GABA\textsubscript{A} receptors that mediate tonic inhibition\textsuperscript{35}. Given our results with GABA\textsubscript{A} receptors containing \(\alpha\)4 and \(\alpha\)1 subunits together with the \(\gamma2\) or \(\delta\) subunit, one could assume that DCUK-OEt would well affect the function of synaptically localized GABA\textsubscript{A} receptors as well as certain extrasynaptic GABA\textsubscript{A} receptors. We noted two characteristics of DCUK-OEt that suggest that its primary effect may be at extrasynaptic receptors containing either \(\gamma2\) or \(\delta\) subunit together with an \(\alpha1\) and \(\gamma3\) subunit. These combinations of subunits (\(\alpha\)1\beta3\gamma2 and \(\alpha\)1\beta3\gamma3) display a low EC\textsubscript{50} for GABA (see Supplementary Fig. S6) and DCUK-OEt can produce highly significant potentiation of \(\alpha\)1\beta3\gamma2 and \(\alpha\)1\beta3\gamma3-mediated currents at the EC\textsubscript{10} concentration of GABA in our assays, and probably at concentrations of GABA consistent with those encountered in locations outside of the GABA synapse. This observation would be quite compatible with significant potentiation at extrasynaptic sites where concentrations of GABA have been considered to be in the high mM range, as opposed to the high concentrations (mM) of GABA that are present in the synapse\textsuperscript{48}. We saw no measurable effect of DCUK-OEt on \(\alpha\)1\beta3\gamma2 receptors at high concentrations of GABA (EC\textsubscript{40} and above), and non-significant effects on \(\alpha\)1\beta3\gamma2 and \(\alpha\)1\beta3\gamma1 GABA\textsubscript{A} receptors at low GABA concentrations (EC\textsubscript{40}). Since \(\alpha\)1\gamma3 is responsible for the major portion of the phasic actions of GABA, and relatively high amounts of \(\beta1\) and \(\gamma1\) were reported at synaptic sites in CeA\textsuperscript{10,32-34}, it is plausible that phasic effects of GABA through these subunit combinations would not be modulated by DCUK-OEt. In fact, when we applied DCUK-OEt focally to CeA neurons, we found no change in sPSC frequency, amplitude, rise or decay time, indicating no effects on phasic transmission (Fig. 4c).

There is strong evidence for the existence of \(\alpha\)1\beta3\delta receptors located extrasynaptically in particular areas of brain (i.e., the interneurons of the hippocampus and particularly those of the dentate gyrus)\textsuperscript{33-35}. Tonic inhibition mediated by GABA\textsubscript{A} receptors containing the \(\alpha1\) subunit has also been noted in the CeA\textsuperscript{36}. Our prior studies using slices of the CeA demonstrated that CRF1 receptor-positive (CRF1+) neurons express the \(\alpha1\) GABA\textsubscript{A} receptor subunit, and this subunit is integral for the GABA-mediated tonic conductance in these neurons as well as being involved in the phasic synaptic response to GABA\textsuperscript{36}. When we measured tonic conductance in CeA neurons, focal application of DCUK-OEt produced an enhancement of the recorded tonic current, suggesting local effects of DCUK-OEt at extrasynaptic GABA\textsubscript{A} receptors. To further ascertain whether the effects of DCUK-OEt were mediated particularly by extrasynaptic GABA\textsubscript{A} receptors, we performed a comparison of the change (increase) in current produced by DCUK-OEt and the decrease generated by the subsequent co-application of 100 \(\mu\)M gabazine\textsuperscript{37}. The strong correlation indicated that DCUK-OEt was indeed stimulating
a tonic conductance in these neurons by actions at extrasynaptic GABA_A receptors. Recently, de Guglielmo et al.38 reported that inactivation of an ensemble of neurons in the CeA resulted in abrogation of excessive alcohol consumption by alcohol-dependent rats. The anatomical area of the CeA from which we obtained our electrophysiologic data coincides with the area containing the ensemble described by de Guglielmo et al.38 An increase in the tonic conductance through extrasynaptic GABA_A receptors, mediated by DCUK-OEt, may engender reduced activity of the neurons identified by de Guglielmo et al.38 and be an effective mode for reducing alcohol intake by dependent animals.

In all, our characterization of DCUK-OEt indicates that this molecule has characteristics that resemble those of etomidate, other anesthetics, ethanol and neurosteroids, but the full profile of DCUK-OEt actions speaks to an interaction with a site or sites on the GABA_A receptor that distinguish DCUK-OEt from currently known PAMs and direct agonists acting at GABA receptors.

Methods

Radioligand binding. [3H]Flunitrazepam Binding and Displacement by DCUK-OEt. Membrane Preparation. These experiments were performed at the University of Colorado Health Sciences Center, Denver, CO. Experiments were approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Colorado, Denver, and were performed in accordance with the NIH Guide for the Care and Use of Laboratory Animals. Male Sprague-Dawley rats (200–250 g) were maintained in an AAALAC-accredited facility and sacrificed by CO2 exposure and decapitation. Brains were removed, and membranes were prepared from the forebrain of Colorado, Denver, and were performed in accordance with the NIH Guide for the Care and Use of Laboratory Animals. The complementary DNAs encoding the GABA_A subunits were synthesized de novo, optimized for Xenopus laevis oocyte expression and subcloned in pGEMHE by GenScript (Piscataway, NJ). The in vitro transcription of GABA_A subunits was performed using mMessage mMachine (Life Technologies, Grand Island, NY). After isolation of Xenopus laevis oocytes, they were injected with capped complementary RNAs encoding wild-type or mutant subunits in different ratios, depending on the subunits: α1/α2:2, 2:2:20 ng; α1/β1/γ2:1, 2:2:6 ng; α1/β2/γ3:3:3 ng; α3/β3/γ3:3, 0.5:0.5:5 ng; α3/β3/γ4:3:3:2, 0.1:0.1:1 ng; α1/γ3:3, 0.5:0.5 ng; α1/β3:3:3:3, 0.4:0.4:4 ng.

Electrophysiology. Oocyte electrophysiology. Xenopus laevis frogs were obtained from Nasco (Fort Atkinson, WI, USA). All surgery was performed in accordance with a protocol approved by the University of Texas, Austin IACUC and the NIH Guide for the Care and Use of Laboratory Animals. The complementary DNAs encoding the GABA_A subunits rat α1, β1, γ1, and human β2 were provided by Drs Myles H. Akabas, Paul J. Whiting and Richard W. Olsen. Human γ1 cDNA was synthesized de novo, optimized for Xenopus laevis oocyte expression and subcloned in pGEMHE by GenScript (Piscataway, NJ). The in vitro transcription of GABA_A subunits was performed using mMessage mMachine (Life Technologies, Grand Island, NY). After isolation of Xenopus laevis oocytes, they were injected with capped complementary RNAs encoding wild-type or mutant subunits in different ratios, depending on the subunits: α1/β2:2, 2:2:20 ng; α1/β2/γ1:1, 2:2:6 ng; α1/β2/γ3:3:3 ng; α3/β3/γ3:3, 0.5:0.5:5 ng; α3/β3/γ4:3:3:2, 0.1:0.1:1 ng; α1/γ3:3, 0.5:0.5 ng; α1/β3:3:3:3, 0.4:0.4:4 ng.

Electrophysiology. The injected oocytes were incubated at 15°C in sterilized Modified Barth’s solution for 1–7 days before recording, and the responses of GABA_A receptors expressed in oocytes were studied using two-electrode voltage clamp as described earlier.33,66 Oocytes were discarded if the maximal current was over 20 µA or if the baseline was unstable or drifted to positive values.
Recording protocols. **GABA concentration-response curves.** Increasing concentrations of GABA were applied for 20–30 s (0.1–1000 μM) followed by 5–15 minutes washout. Responses were expressed as percentages of the maximal current (Supplementary Fig. S6).

**Modulator application.** DCUK-OEt and DCUKA stock solutions were prepared in DMSO weekly, then sonicated for 5 min, and stored at 4 °C, protected from light. On the day of the experiment, dilutions were prepared, sonicated for 5 min, and used immediately. The final DMSO concentration in the buffer bathing the oocyte was ≤0.1%. In order to test the effects of DCUKs, the agents were first pre-applied for 1 min and then co-applied with GABA. To verify the presence of a third subunit in expressed subunit combinations, the responses to GABA in the presence of Zn++ (1, 10 or 100 μM) were evaluated (Supplementary Table S2). The application sequence in each instance was as follows: Maximal GABA (20 s application, 15 min washout), EC10 GABA (30 s application, 5 min washout), EC10 GABA, pre-application of the drug followed by a co-application with EC10 GABA, EC10 GABA, pre-application of Zn++ immediately followed by a co-application with EC10 GABA, EC10 GABA. In most cases, we limited to one DCUK application per oocyte. Flumazenil and 17PA were pre-applied before their co-application with GABA. When co-applying with DCUK, the antagonist and DCUK were pre-applied together before their co-application with GABA. FluNitrazepam was not pre-applied before co-application with GABA.

**Statistical Analysis.** Responses to DCUK-OEt were quantified as the percent change in current between the response to the EC90 concentration of GABA and the response to the EC10 concentration of GABA in the presence of 0.3 μM concentration of DCUK-OEt. To control for batch effects a linear mixed model was implemented in SAS (version 9.4) to calculate the normalized percent change in current for each receptor subunit combination (each receptor combination was examined in two to nineteen separate experiments). Because each receptor was examined across several experiments, a random effect of batch was included in the model. For each receptor, the estimated percent change in the GABA EC10-induced current produced by addition of DCUK-OEt was compared to zero by ascertaining whether zero percent change was outside the confidence interval of the measured values. This was accomplished by using a single sample t-test in the MIXED procedure of SAS, and a Bonferroni adjustment to control for multiple comparisons across receptors. Comparisons between receptors with a single subunit difference were executed within the linear mixed model using linear contrasts. A Bonferroni adjustment was used to control for multiple pairwise comparisons. Significant effects are those with a Bonferroni adjusted p-value < 0.05 and marginal effects are those with a Bonferroni adjusted p-value < 0.2.

Other statistical tests (t-test and ANOVA) were applied as indicated in the corresponding table or figure legend.

The GABA concentration response curves (CRCs) were fitted to the following equation:

$$I/I_{\text{MAX}} = \frac{1}{1 + 10^{\log EC_{50} - \log (GABA)} \times n_H}$$

where $I/I_{\text{MAX}}$ is the fraction of the maximally-obtained GABA response, $EC_{50}$ (effective concentration 50) is the concentration of GABA producing a half-maximal response, [GABA] is GABA concentration and $n_H$ is the Hill coefficient.

**Brain slice electrophysiology.** **Brain slice preparation.** All procedures were approved by the Scripps Research IACUC and were consistent with the NIH Guide for the Care and Use of Laboratory Animals. Slices were prepared from brains of 5 adult male Wistar rats (250–350 g) as described by Herman et al.88. A single slice was transferred to a recording chamber mounted on the stage of an upright microscope (Olympus BX50W1).

**Brain slice electrophysiological recording.** Neurons were visualized and whole cell patch clamp recordings were made as previously described88. Series resistance was typically <15 MΩ and was continuously monitored with a hyperpolarizing 10 mV pulse. Electrophysiological properties of cells were determined by pClamp 10 Clampex software online during voltage-clamp recording using a 5 mV pulse delivered after breaking into the cell. The resting membrane potential was determined online after breaking into the cell using the zero current (I = 0) recording configuration and the liquid junction potential was included in the determination.

DCUKA and DCUK-OEt were prepared as described for the experiments on oocyte electrophysiology. Other drugs were dissolved in aCSF, and all drugs were applied by Y-tubing application for local perfusion primarily on the neuron of interest. To isolate the inhibitory currents mediated by GABA$_A$ receptors, all recordings were performed in the presence of glutamate and GABA$_A$ receptor blockers88. All voltage clamp recordings were performed in a gap-free acquisition mode with a sampling rate per signal of 10kHz or a total data throughput equal to 20kHz (2.29 MB/min) as defined by pClamp 10 Clampex software.

**Data Analysis.** Frequency, amplitude and decay of spontaneous inhibitory postsynaptic currents (sIPSCs) were analyzed and visually confirmed using a semi-automated threshold based mini detection software (Mini Analysis, Synaptosoft Inc.). Averages of sIPSC characteristics were determined from baseline and experimental drug conditions containing a minimum of 60 events (time period of analysis varied as a product of individual event frequency) and decay kinetics were determined using exponential curve fittings and reported as decay time (ms). All detected events were used for event frequency analysis, but superimposed events were eliminated for amplitude and decay kinetic analysis. In voltage clamp recordings, tonic currents were determined using Clampfit 10.2 (Molecular Devices) and a previously-described method88. Responses were quantified as the difference in holding current between baseline and experimental conditions. Events were analyzed for independent significance using a one-sample t-test and compared using a two-tailed t-test for independent samples, a paired two-tailed t-test for comparisons made within the same recording, and a one-way ANOVA with a Bonferroni post
Molecular modeling. All molecular modeling studies were conducted using Biovia Discovery Studio 2016 (Biovia Inc., San Diego, CA) and all crystal structure coordinates were downloaded from the protein data bank (www.pdb.org). The homology model of the human GABA<sub>A</sub> receptor pentamer was generated with the MODELLER protocol<sup>82</sup> using the crystal structures of the human GABA<sub>A</sub> receptor<sub>33</sub> homopentamer as a template (PDB ID: 4COF<sup>3</sup>, Uniprot accession: P14867) and 2<sub>2</sub> (Uniprot accession: P18507) subunits were superimposed over the template, with the crystal structure of two 33 subunits, so that the final pentameric model consisted of two α<sub>1</sub>, two 33, and one 2<sub>2</sub> subunits, arranged in an αβ3β3α<sub>2</sub> pattern (counterclockwise, as seen from above). The resulting final structures were subjected to energy minimization utilizing the conjugate gradient minimization protocol with a CHARMM forcefield and the Generalized Born implicit solvent model with simple switching (GBSW)<sup>64</sup>. The minimization calculations converged to a RMS gradient of < 0.01 kcal/mol. The Flexible Docking protocol<sup>65</sup>, which allows flexibility in both the protein and the ligand during the docking calculations, was used to predict the binding orientations of both known and candidate GABA<sub>A</sub> PAMs in the binding site located at either the classical α<sub>1</sub>-γ<sub>2</sub> benzodiazepine site (α + γ<sub>2</sub>-interface) or the alternative α<sub>1</sub>-β<sub>2</sub> site (α + β<sub>2</sub>-interface). Predicted binding poses were energy-minimized in situ using the CDocker protocol<sup>66</sup> prior to calculation of binding energies using the distance-dependent dielectric model. All numeration refers to the corresponding mature protein.

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**Author Contributions**

C.M.B.: Performed the electrophysiologic studies with oocytes, analyzed part of the data and participated in writing the manuscript. M.H.: Performed the electrophysiologic studies with brain slices and assisted in writing the manuscript. L.D.S.: Performed the receptor binding studies and analyzed that data. K.J.L.: Assisted in oocyte electrophysiology and collected data. H.-Y.L.: Assisted in oocyte electrophysiology and collected data. D.S.B.: Performed the docking experiments, analyzed the computational results and assisted in writing the manuscript. L.A.V.: Performed the statistical analysis of part of the oocyte electrophysiology data. R.A.H.: Established the experiments with oocytes, reviewed all data and advised on writing the manuscript. M.R.: Supervised the electrophysiology studies with brain slices and assisted in writing the manuscript. P.L.H.: Supervised all of the receptor binding studies and helped write and edited the manuscript. B.T.: Generated the diphenylureido quinoline compounds used in these studies, conceived the experimental studies, coordinated experiments, interpreted data, wrote the manuscript.

**Additional Information**

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**Competing Interests:** DCUKA and DCUK-OEt are patented products of Lohocla Research Corporation (U.S. Patent #s 6,962,930 and 7,923,458 and PCT Application No. PCT/US2015/036473). Boris Tabakoff is CEO and CSO of Lohocla Research Corporation. Paula Hoffman is an employee and member of the Scientific Advisory Board of Lohocla Research Corporation. Lawrence Snell was a previous employee of Lohocla Research Corporation. The other authors declare no competing financial interest to this work.

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