PHARMACOLOGY AND CELL METABOLISM

Tryptophan in Alcoholism Treatment I: Kynurenine Metabolites Inhibit the Rat Liver Mitochondrial Low \( K_m \) Aldehyde Dehydrogenase Activity, Elevate Blood Acetaldehyde Concentration and Induce Aversion to Alcohol

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Abstract — Aims: The aims were to provide proofs of mechanism and principle by establishing the ability of kynurenine metabolites to inhibit the liver mitochondrial low \( K_m \) aldehyde dehydrogenase (ALDH) activity after administration and in vivo, and to induce aversion to alcohol. Methods: Kynurenic acid (KA), 3-hydroxykynurenine (3-HK) and 3-hydroxyanthranilic acid (3-HAA) were administered to normal male Wistar rats and ALDH activity was determined both in vitro in liver homogenates and in vivo (by measuring blood acetaldehyde following ethanol administration). Alcohol consumption was studied in an aversion model in rats and in alcohol-preferring C57 mice. Results: ALDH activity was significantly inhibited by all three metabolites by doses as small as 1 mg/kg body wt. Blood acetaldehyde accumulation after ethanol administration was strongly elevated by KA and 3-HK and to a lesser extent by 3-HAA. All three metabolites induced aversion to alcohol in rats and decreased alcohol preference in mice. Conclusions: The above kynurenine metabolites of tryptophan induce aversion to alcohol by inhibiting ALDH activity. An intellectual property covering the use of 3-HK and 3-HAA and derivatives thereof in the treatment of alcoholism by aversion awaits further development.

INTRODUCTION

It is generally accepted that the most effective relapse preventing drugs following detoxification of alcohol-dependent subjects are the \( N \)-methyl-D-aspartate modulator acamprosate, the \( \mu \)-opioid receptor antagonist naltrexone and the aldehyde dehydrogenase (ALDH) inhibitor and alcohol aversion drug disulfiram (Mann, 2004). The superiority of disulfiram over acamprosate and naltrexone has been demonstrated (see, e.g. de Sousa and de Sousa, 2004, 2005; Laaksonen et al., 2008; Diehl et al., 2010). However, safety issues with disulfiram (Chick, 1999) suggest the need for developing safer alternative ALDH inhibitors for alcoholism treatment. One such alternative may be the essential amino acid \( L \)-tryptophan (Trp) and its metabolites.

Of all amino acids, Trp has been the most extensively studied in relation to alcohol and alcoholism. While Trp metabolism and disposition are greatly influenced by acute and chronic alcohol (ethanol) consumption and subsequent withdrawal in both humans and experimental animals (for reviews, see Le Marquand et al., 1994a, b; Badawy, 2002, 2005), Trp itself can also influence alcohol consumption. Thus, alcohol consumption by rats is decreased by Trp and its 5-hydroxylated metabolite, although the Trp effect is controversial (for review and references, see Naranjo et al., 1986). The implication of serotonin (5-hydroxytryptamine or 5-HT) as a modulator of alcohol consumption has arisen from a variety of studies using treatments influencing the metabolism and function of this indolylamine, such as its Trp and 5-hydroxytryptophan precursors, the Trp hydroxylase inhibitor \( p \)-chlorophenylalanine, various serotonin reuptake inhibitors, serotonin postsynaptic receptor activators and neurotoxic destruction of serotonin neurons, reviewed by Naranjo et al., 1986; Sellers et al., 1992). Accordingly, it has always been assumed that the decrease in alcohol consumption by Trp is mediated by serotonin, although no mechanistic studies were performed to confirm this assumption. An alternative or additional mechanism is that of Trp acting peripherally via its metabolites to induce aversion. Thus, we have previously reported (Badawy and Morgan, 2007) that a number of Trp metabolites of the quantitatively most important of the Trp-degradative pathways (the hepatic kynurenine pathway; Fig. 1), namely 3-hydroxykynurenine (3-HK), 3-hydroxyanthranilic acid (3-HAA) and kynurenic acid (KA), and also the Trp transamination product indol-3-ylpyruvic acid (IPA) strongly inhibit the activity of the mammalian (rat) liver mitochondrial low \( K_m \) ALDH in vitro. At 2 \( \mu \)M, inhibition by these four metabolites was 55, 46, 40 and 30% respectively, against a 42% inhibition by a similar concentration of the classical ALDH inhibitor disulfiram. Less strong inhibition was also observed with the kynurenic metabolite xanthurenic acid and with 5-hydroxytryptophan and it is of interest that inhibition of alcohol consumption by this immediate serotonin precursor has been suggested to involve a peripheral, in addition to a central and component (Zabik et al., 1994). In the present paper, we demonstrate the ability of the former three kynurenine metabolites to: (a) inhibit ALDH activity after administration to rats; (b) elevate blood acetaldehyde concentration following acute ethanol administration; (c) induce aversion to alcohol in an experimental aversion model and (d) decrease alcohol preference in mice, and propose these metabolites as potential alcohol aversion drugs. The Trp transamination metabolite IPA was not included in the present work because of its back-conversion into Trp (Richards et al., 1972).
MATERIALS AND METHODS

Chemicals and other materials
Bovine albumin, disulfiram (tetraethyl thiuram disulfide), ethanol (>99%), oxidized nicotinamide-adenine dinucleotide (NAD)+, Trp, 3-HK, 3-HAA, KA and other kynurenine metabolites were purchased from the Sigma-Aldrich Co Ltd. (Fancy Road, Poole, Dorset, UK) and were stored as directed by the manufacturer. Water and methanol [high-performance liquid-chromatographic (HPLC) grade] were purchased from either VWR International (Hunter Boulevard, Magna Park, Leicestershire, UK) or Fisher Scientific UK (Bishop Meadow Road, Loughborough, Leicestershire, UK). Acids and alkalis were purchased from VWR International, were of the purest commercially available grades and were made up in HPLC-grade water. Filtration, Eppendorf and other tubes were purchased from Fisher or other standard suppliers.

Animals and treatments
Adult normal male Wistar rats weighing between 150 and 170 g at the start of experiments were purchased from accredited animal suppliers and were acclimatized to our standard UK Home Office-approved housing conditions (21 ± 2°C, relative humidity of 55 ± 10% and a 12 h/12 h light: dark cycle) for at least one week before experiments. They were housed five per cage, unless stated otherwise, in conventional open-top cages with standard softwood bedding from accredited suppliers, and were allowed free access to standard laboratory RM1 diet and water. Adult male mice of the alcohol-preferring C57BL/6J strain, weighing between 20 and 22 g at the start of experiments were also investigated. This study was performed under the auspices of Cardiff University and approved and licensed (project licence No: PPL 30/2502) by the UK Home Office under the Animal (Scientific Procedures) Act 1986. All compounds were administered intraperitoneally in 0.9% (w/v) NaCl (physiological saline) or, in the case of disulfiram, in a mixture of dimethylformamide: saline (1:1). 3-HK, 3-HAA and KA were given in single doses of 1–10 mg/kg body wt, whereas disulfiram was given in a 100 mg/kg dose. When given repeatedly for up to 8 days, these compounds were given in the above single doses once daily.

Determination of ALDH activity
The mitochondrial low K<sub>m</sub> ALDH activity was determined by the method of Tottmar <i>et al.</i> (1973) in rat liver supernatants prepared as described by Mazzanti <i>et al.</i> (1989). Briefly, a 1 g piece of frozen liver was homogenized in 9 ml of an ice-cold homogenization buffer consisting of 0.25 M sucrose, 5 mM Tris–HCl and 0.5 mM ethylene-diamine tetra-acetic acid disodium salt (pH 7.2) for 1 min using an ultra-Turrax homogenizer. The homogenate was centrifuged at 500 g for 15 min at 4°C. The decanted supernatant was treated with 0.4 ml of a 5% aqueous sodium deoxycholate solution, stirred gently and made up to 10 ml with the above homogenization buffer. For assay of the low K<sub>m</sub> ALDH activity, the 1 ml total incubation mixture contained the following components in their final concentrations: NAD+ (1 mM), pyrazole (0.1 mM), rotenone (2 μM), sodium pyrophosphate buffer, pH 8.8 (50 mM), acetaldehyde (5 μM) and 0.1 ml of the above liver supernatant. Before the addition of acetaldehyde, the mixture was preincubated for 10 min at 25°C. The reaction was then started by the addition of acetaldehyde followed by incubation at 25°C for 5 min with shaking, and was terminated by placing the incubation tubes on ice. The reduced nicotinamide-adenine dinucleotide formed was determined by measuring its absorption at 340 nm against a blank preincubated and incubated as above, but to which water was added instead of acetaldehyde. Blanks were performed in duplicates, whereas tests were in triplicates. ALDH activity was expressed in μmol of NADH formed/min per mg of protein. The latter was determined by the method of Lowry <i>et al.</i> (1951), using bovine serum albumin as standard. In initial experiments, the high K<sub>m</sub> ALDH activity was also determined in the presence of a 5 mM acetaldehyde concentration.

Determination of ALDH activity in vivo
ALDH activity in vivo was determined by measuring blood acetaldehyde concentration following acute ethanol administration. The recovery of acetaldehyde from rat blood is best achieved by haemolysis of the blood sample, rather than by acid precipitation (Eriksson <i>et al.</i>, 1977; Eriksson, 1980). Rats received an intraperitoneal injection of kynurenine metabolites (10 mg/kg body wt each). One hour later, another injection of ethanol (2 g/kg body wt as a 25% v/v solution in physiological saline) was administered. Blood samples (100 μl each) were withdrawn at hourly intervals for...
3 h after the ethanol injection from a tail vein under light iso-
fluorane anaesthesia and immediately added to 0.9 ml of
ice-cold water in a gas-chromatographic (GC) injection vial.
The vial was immediately sealed using a crimper and the
contents of the vial were mixed. The vials were subsequently
stored in a refrigerator at 4°C overnight before analysis the
following morning. Acetaldehyde and ethanol in the sealed
vials were analyzed by head-space GC using a Perkin Elmer
Clarus 500 gas chromatograph with a Turbo-Mix HS 40
autosampler. GC conditions were as follows: injection
needle temperature: 85°C; transfer temperature: 70°C; oven
temperature: 80°C; pressure: 13 psi; sample equilibration
time: 22.3 min at 65°C. The system was operated by the
associated Total Chrome software, which also controlled data
handling and processing. A 100 μl portion of a standard
mixture of ethanol and acetaldehyde (1 mg/ml each), pre-
pared in the Laboratory by diluting suitable amounts of
ethanol and acetaldehyde previously stored at 4°C with cold
deionized water was diluted with 0.9 ml of cold water and
run as calibrant at the start of each analytical run. Initially,
this standard mixture was calibrated against a certified foren-
sic analytical 6-component standard containing ethanol and
acetaldehyde at 1 mg/ml each (Restek).

Pitfalls in acetaldehyde determination

One such pitfall is artifactual formation of acetaldehyde
from ethanol in the presence of blood, by a process depend-
et on ethanol but not on its concentration (Truitt, 1970;
Eriksson et al., 1977; Eriksson, 1980). This was confirmed
in the present work in a control experiment in which
ethanol was added in final concentrations of 10–100 mM
to 0.1 ml portions of normal untreated rat blood. The
mixture was diluted to 1 ml with water in GC vials, which
were then sealed and analyzed as above. Acetaldehyde was
formed in amounts (in μM) of 61, 63, 64, 69, 61, 65 and
73 when ethanol was present at concentrations of 10, 20,
30, 40, 50, 75 and 100 mM, respectively. The acetaldehyde
formed was subtracted from the in vivo experimental
values observed.

Another problem in acetaldehyde determination is its
binding to haemoglobin and the consequently lower recov-
er. The above authors recommended that, for rat blood, this
problem can be largely overcome by haemolysis of blood
samples (by dilution of one part of blood with nine parts of
water) prior to head-space GC analysis and by performing
control experiments to assess acetaldehyde recovery.
Acetaldehyde recovery from rat blood was reported by the
above authors to be 90% when acetaldehyde was added at a
final concentration of 100 μM. This was also confirmed in
the present study, in which a recovery value (expressed as
the mean percentage recovery ± SEM for three determina-
tions) of 90.5 ± 3.8 was observed with the above acet-
aldehyde concentration. Recoveries at other concentrations
of added acetaldehyde (25, 50, 150 and 200 μM) were 91.5 ±
2.8, 80.4 ± 2.7, 84.5 ± 2.4 and 85.4 ± 1.9%, respectively. The
blood acetaldehyde concentration values in Fig. 2b were,
however, not corrected for full recovery because of the sim-
ultaneous presence of ethanol under the experimental con-
ditions of Fig. 2. Eriksson et al. (1977) found that, in the
presence of ethanol, the 90% recovery value rose to 105.4%
and it was therefore considered unnecessary to apply a
recovery factor, which in any case would have little effect on
the results in Fig. 3b.

We have thus addressed the above pitfalls in detail, taken
appropriate steps to circumvent them and are therefore confi-
dent of the validity of the blood acetaldehyde values reported
in the present study.

Determination of liver Trp and kynurenine metabolite
centralations

Trp and its kynurenine metabolites were determined in liver by
our newly developed rapid isocratic high-performance liquid-
chromatographic procedure (Badawy and Morgan, 2010).
Briefly, a Perkin Elmer LC200 system consisting of a qua-
ternary pump, a column oven and a degasser was used with ultra-
violet and fluorimetric detection in series. The mobile phase
was a methanol: sodium dihydrogen phosphate mixture (27:73,
by vol) at a final pH of 2.0 or 2.8. The system was run isocrati-
cally using a Synergi 4 μ reverse-phase Fusion-RP80 A
column (250 × 4.6 mm) with guard column (Phenomenex).
Operation of the system, data processing and handling were all
performed by the associated Total Chrome software. A standard mixture of Trp and six of its kynurenine metabolites (1 μg/ml each) was used as calibrant at the start of each run. Results were corrected for full recovery.

Alcohol aversion test

Aversion to alcohol was assessed using the alcohol aversion model of Garver et al. (2000). Briefly, individually housed rats were allowed free access to food and water for 3 days. Disulfiram (100 mg/kg), kynurenine metabolites (10 mg/kg each) or vehicle were injected intraperitoneally once daily for 4 days. On the evening of the third day, water (but not food) was withdrawn for 18 h, but was then replaced by a 6% (v/v) ethanol solution, 2 h after the injection on the fourth day. Body wt was measured daily and alcohol consumption was monitored hourly for 4 h and levels were expressed cumulatively over this duration in g/kg body wt.

Alcohol consumption and preference in alcohol-preferring C57BL/6J mice

The potential effects of kynurenine metabolites on alcohol consumption and preference by male alcohol-preferring C57BL/6J mice were also studied. No attempt was made to enhance alcohol preference by acclimatizing the mice to increasing ethanol concentrations, as the purpose of our study was to investigate aversion, which is the proposed primary mechanism of action of kynurenine metabolites, rather than preference. Accordingly, four groups (n = 8 each) of individually housed mice were given free choice of drinking water and a 10% (v/v) ethanol solution for 3 weeks to establish their drinking patterns. Thereafter, mice received a single daily intraperitoneal injection of kynurenine metabolites (10 mg/kg body wt each) or an equal volume of saline for 5 days. Body wt and water and ethanol consumption were recorded daily throughout the whole study duration. Determined per kg body wt, daily alcohol consumption was expressed in absolute amounts (g) and as a percentage of total fluid intakes (% preference).

Statistical analysis

Enzymatic and other biochemical test results were compared with those of control groups by the unpaired t test, whereas alcohol consumption results were assessed initially by one-way analysis of variance (ANOVA) and additionally for within-group differences (time factor versus baseline values) by paired t-tests, using Sigma Plot (Systat, UK), version 11, with which graphics were prepared. For multiple group comparisons using this program, the Holm-Sidak test was applied, as it is more powerful than the Tukey or Bonferroni tests and can be used for both pairwise comparisons and those versus a control group. Where the data failed the normality (Shapiro–Wilk) test, Kruskal–Wallis one-way ANOVA on ranks was performed. A two-tailed level of significance (P) was set at 0.05.

RESULTS

Inhibition of the rat liver mitochondrial low K_m ALDH activity by acute and chronic administration of kynurenine metabolites

Acute time-course (Fig. 2a) and dose–response (Fig. 2b) experiments were performed. Initially, we found that the high K_m ALDH activity (assayed in the presence of 5 mM acetaldehyde) was not influenced by kynurenine metabolites. Consequently all results reported in this section concern the low K_m activity. As shown in Fig. 2a, the mitochondrial low K_m ALDH activity was inhibited by all three kynurenine metabolites after administration of a 10 mg/kg body wt dose. At 0.5 h after administration, ALDH activity was inhibited by 3-HK and KA by 45 and 37%, respectively (P = 0.0267–0.0024). Inhibition by all three metabolites was then maintained at 40–51% at 1 h. Thereafter, inhibition by KA remained at this latter level until 4 h, whereas that by 3-HAA continued to strengthen, reaching 84% at 4 h (P = 0.0000). With 3-HK, ALDH activity began to recover at 2 h, but remained significantly inhibited at 2 and 3 h (by 30 and 17%, respectively; P = 0.0128–0.0431).
As shown in Fig. 2b, significant inhibition of 32–56% \((P = 0.0398–0.0008)\) of ALDH activity was observed at 1 h after administration of a 1 mg/kg body wt dose of kynurenine metabolites. Maximum inhibition at 1 h was observed with a 2.5 mg/kg dose of KA (86%) and with a 7.5 mg/kg dose of 3-HK (59%) and 3-HAA (70%) \((P = 0.0035–0.0001)\).

ALDH activity was also inhibited when a 10 mg/kg body wt dose of the above 3 kynurenine metabolites was administered daily for 8 days (data not shown). At 2 h after the (final) injection on the 8th day, inhibition by KA, 3-HK and 3-HAA was 37%, 39% and 64% respectively \((P = 0.0292–0.0007)\). Although this inhibition could very well be due to the acute effect of kynurenine metabolites, it suggests that no tolerance develops towards it after chronic treatment.

**Inhibition of ALDH activity in vivo by acute administration of kynurenine metabolites**

ALDH activity in vivo was determined by measuring the accumulation of acetaldehyde in blood following acute ethanol administration. The results in Fig. 3 show blood-ethanol (a) and acetaldehyde (b) concentrations after intraperitoneal administration of a 2 g/kg body wt dose of ethanol. In saline-pretreated control rats, ethanol concentration rose to 35.8 mM at 1 h and to 37.1 mM at 2 h before declining to 26.0 mM at 3 h. None of the three kynurenine metabolites exerted a significant effect on ethanol concentration at 1 h \((P > 0.1)\). With 3-HAA, ethanol concentration resembled that in saline-treated controls at 2 and 3 h. By contrast, ethanol concentration at 2 and 3 h after ethanol administration was significantly decreased by pretreatment of rats with KA and 3-HK, by 24–27% \((P = 0.05–0.0175)\).

Blood acetaldehyde concentration following ethanol administration (Fig. 3b) to saline-pretreated control rats remained at a constant level of 45–47 \(\mu\)M, suggesting a constant rate of ethanol and acetaldehyde oxidation over the 1–3 h observation period. Pretreatment of rats with KA increased acetaldehyde concentration by 109, 125 and 115%, respectively \((P = 0.0104–0.0029)\). 3-HK induced a stronger elevation of acetaldehyde concentration, of 180, 174 and 202%, respectively at 1–3 h after ethanol administration \((P = 0.0017–0.0000)\), whereas with 3-HAA, the elevation of acetaldehyde concentration was modest (42, 34 and 56%, respectively), and not significant \((P > 0.1)\). The elevation of blood acetaldehyde concentration by 3-HK was significantly greater than that by KA at 1–3 h \((P = 0.05–0.0095)\).

**Demonstration of aversion to alcohol after administration of kynurenine metabolites and disulfiram**

In the aversion model of Garver et al. (2000), rats treated with the classical ALDH inhibitor disulfiram consumed equal amounts of the ethanol drinking solution as control animals during the first hour of the test. Thereafter, alcohol consumption by disulfiram-treated rats remained static, unlike that by controls, which continued to increase cumulatively up to the fifth hour. The results in Fig. 4a, show that disulfiram actually inhibited alcohol consumption significantly and maximally during the first hour (by 51%; \(P = 0.0351\), paired t-test), thereafter the animals continued to drink the ethanol solution, but to a lesser degree than controls. Thus, the inhibition of ethanol consumption by disulfiram was maintained for two more hours, as 45–46% \((P = 0.0158–0.0138)\), but, at 4 h, the 37% decrease was not significant. The results in Fig. 4b show that kynurenine metabolites also inhibited alcohol consumption in this model significantly \((P = 0.05–0.005)\) over the first 3 h. Thus, as was the case with disulfiram, inhibition of alcohol consumption was strongest at 1 h after administration of KA, 3-HK and 3-HAA (by 28, 26 and 50%, respectively). Inhibition remained significant at 2 and 3 h, but, by 4 h, only that by KA was still significant.

It will be noted from the data in Fig. 4 that alcohol consumption by control rats in the disulfiram experiment (Fig. 4a) is lower than that of the control animals in the kynurenine metabolite experiment (Fig. 4b). This is almost certain to be due to the use of dimethylformamide, along with saline, to dissolve disulfiram, rather than to variations among different batches of animals, because, as will be seen in the accompanying paper (Badawy et al., 2011), the control data obtained in animals given saline only in two different experiments were broadly similar to those in the present experiment with kynurenines.
Alcohol consumption by C57 mice

As no attempt was made to enhance alcohol preference by acclimatizing the mice to increasing ethanol concentrations, there were wide variations in levels of consumption of the 10% (v/v) ethanol solution, with ~37–50% of the mice in each group consuming >10 g/kg/day, and showing >50% preference, with the remainder consuming up to 7 g/kg/day with a level of preference <32%. To establish comparability between groups at baseline (Day 0), six mice from each group were selected whose results are shown in Fig. 5. As shown, the % preference (Fig. 5a) and absolute ethanol intake in g/kg body wt (Fig. 5b) were broadly similar across groups at baseline, with no significant differences ($P > 0.1$). In the saline-treated control group, preference remained stable for 4 days and the decrease on Day 5 did not reach statistical significance ($P = 0.095$). By contrast, preference decreased in mice receiving the three kynurenine metabolites, with 3-HAA causing the largest decrease. Compared with baseline, the decrease in the % preference with 3-HAA (23–46%) was significant on all days ($P = 0.043–0.009$), except Day 4. With 3-HK and KA, only the decreases on Days 4 and/or 5 (19–39%) were significant ($P = 0.05–0.011$). When the % preference values with kynurenine metabolites were compared with those of the control group, significant decreases were also observed with 3-HK (37%) on Day 4, with KA (26–30%) on Days 1 and 4 and with 3-HAA (26–36%) on Days 1–5 ($P = 0.05–0.0075$).

When alcohol consumption was expressed in absolute amounts (g/kg body wt) (Fig. 5b), a broadly similar pattern emerged, with 3-HAA causing the greatest decrease in alcohol intake. However, the only significant differences were those compared with the saline controls for KA on Days 1 and 4 and for 3-HAA on Days 1 and 5 (31–45%; $P = 0.021–0.001$).

**Hepatic concentrations of kynurenine metabolites after their administration**

Hepatic concentrations of KA, 3-HK and 3-HAA were determined after acute administration of each individual compound in the same rats in which ALDH activity was assessed. Although concentrations of other kynurenine metabolites and of Trp were also determined, these are not reported here, as their relevance largely falls outside the

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**Fig. 5. Effects of repeated administration of kynurenine metabolites on alcohol consumption and preference in male alcohol-preferring C57BL/6J mice** Experimental details are as described in the ‘Materials and Methods’ section. Alcohol consumption was monitored daily and is expressed (per kg body wt) for each group from Day 0 (before the injections) onwards, both as a preference % (a) and in g amounts (b). Values are means ± SEM (bars) for each group of six mice. For statistical comparisons, see the relevant text in the ‘Results’ section.

**Fig. 6. Hepatic kynurenine metabolite concentrations after their acute administration** Kynurenine metabolite concentrations were determined as described in the ‘Materials and Methods’ section in livers of the same rats undergoing the time-course (a) and dose-response (b) experiments with ALDH reported in Fig. 2. Values are means ± SEM (bars) for each group of five rats. Abbreviations and comparisons are as in Fig. 2.
scope of the present paper. When a 10 mg/kg body wt dose of the three kynurenine metabolites was administered (Fig. 6a), maximum increases (of 4.2–7.3-fold) in their concentrations were observed at 1 h ($P = 0.0002$) and were maintained for 3–4 h. Dose-dependent increases in liver kynurenine metabolite concentrations were observed at 1 h (Fig. 6b) in the 1–10 mg/kg dose-range with KA and 3-HK. However, with 3-HAA, the maximum increase was observed with the 5 mg/kg dose. Baseline [3-HAA] in the time-course experiment was lower than that in the dose–response one. This is most likely due to variations across different animal batches, as the relative increases in this metabolite concentration at 1 h after administration of the 10 mg/kg dose were similar in both experiments (2.67 and 2.97-fold, respectively).

**Animal body weights during repeated administration of kynurenine metabolites and disulfiram**

Body wt was measured in chronic experiments, not only to determine daily dose levels, but also as a measure of animal welfare and safety of administered compounds. In the alcohol aversion study in which compounds were injected once daily for 4 days, both rats treated with kynurenine metabolites and their saline-treated controls gained weight at the normal rate during the first 3 days. However, small losses of 6–9% were observed in both control and test rats on the morning of the final (fourth) day, compared with body weights the day before, almost certainly due to introducing a water-deprivation regimen during the preceding 18 h. In the aversion experiments with disulfiram and its control rats receiving the vehicle (saline: dimethylformamide, 1:1), both control and test rats showed a small wt loss (2.8 and 4.2%) on Day 3, compared with Day 1, presumably due to this solvent, in addition to a 3–6% wt loss following the water-deprivation period, as was the case with kynurenine metabolites and their pure saline control.

In the chronic study of changes in rat liver ALDH activity in which kynurenine metabolites were administered daily for 8 days, changes in body weights were also recorded. As shown in Fig. 7a, all groups gained wt significantly ($P < 0.001$), with the gains by control rats reaching 26% on Day 8. Rats receiving KA and 3-HK also gained wt at a rate close to that of controls (respectively 21 and 24% on Day 8). 3-HAA-treated rats, however, gained wt less strongly, achieving only a 10% wt gain on Days 6–8. All animals appeared to tolerate kynurenine metabolites and showed no adverse reactions or unusual behaviors.

**DISCUSSION**

**Inhibition of ALDH activity after administration of kynurenine metabolites**

The present results have established the ability of three Trp metabolites of the kynurenine pathway, namely KA, 3-HK and 3-HAA to inhibit the rat liver mitochondrial low Km activity of ALDH after acute administration (Fig. 2) and that no tolerance to this inhibition develops after repeated administration. Inhibition was significant with a dose as small as 1 mg/kg body wt (Fig. 2b) and was maximal at 1 h, except after 3-HAA (Fig. 2a). The hepatic concentrations of these metabolites also reached maximum values at 1 h (Fig. 5a) in line with the maximum inhibition. The hepatic concentrations observed (Fig. 5) are well in excess of the 2 μM concentration previously shown to cause 40–55% inhibition in vitro (Badawy and Morgan, 2007).
Inhibition in the present work lasted for at least 3–4 h (the longest durations studied). With 3-HK, ALDH inhibition, which was maximal at 1 h, began to lessen thereafter, suggesting that it is short-lived and hence reversible. However, with KA and 3-HAA, inhibition either remained at its maximal value or gained in strength, suggesting a longer duration and a possible irreversible nature. With disulfiram, its irreversible inhibition (Marchner and Totterm, 1978) is characterized by a prolonged duration (7–10 days) after a single dose (see, Brien and Loomis, 1985). Kinetic studies are clearly required to establish the mechanism(s) of the ALDH inhibition by these kynurenine metabolites.

**Inhibition of ALDH activity in vivo by administration of kynurenine metabolites**

3-HK and KA also inhibited ALDH activity *in vivo*, as suggested by their strong elevation of blood acetaldehyde concentration after ethanol administration (Fig. 3b), whereas 3-HAA caused only a modest increase. As blood-ethanol concentration at 1 h was not influenced by any of the three kynurenine metabolites (Fig. 3a), we conclude that ethanol kinetics were not altered up to this time-point. Thereafter, blood-ethanol concentration was also little altered by 3-HAA, but was significantly decreased by KA and 3-HK, suggesting acceleration of ethanol metabolism by acting on alcohol dehydrogenase or other ethanol-oxidizing enzymes, or through other mechanisms. Further work is required to elucidate the nature of this effect. In relation to a potential acceleration of ethanol metabolism by KA, Lapin and Politi (1994) speculated as to whether the shortening of ethanol-induced sleep time by indol-3-ylpyruvic acid could be due to its conversion to KA. The elevation of blood acetaldehyde concentration by 3-HK and KA strongly suggests that both compounds inhibit ALDH activity *in vivo*, thus further corroborating their effects *in vitro* (Badawy and Morgan, 2007) and after administration (Fig. 2 in the present work).

**Induction of aversion to alcohol by kynurenine metabolites**

The alcohol aversion model of Garver et al. (2000) is a variant of the conditioned taste aversion paradigm in which a novel taste (that of alcohol introduced for the first time) is associated with a noxious condition (the disulfiram-ethanol reaction). That a robust conditioned taste aversion quickly develops under such conditions has been demonstrated (Nolan et al., 1997; Scalera et al., 1997; Barber et al., 1998; Yasoshima and Yamamoto, 1998). We have successfully confirmed the above authors’ findings with disulfiram and demonstrated the ability of our three kynurenine metabolites to induce aversion to alcohol (Fig. 4). Two differences from the findings by Garver et al. (2000) with disulfiram were, however, observed: (a) the inhibition of alcohol consumption by disulfiram was already significant and strongest at the 1 h observation period, whereas the above authors observed no inhibition at 1 h; (b) our disulfiram-treated rats continued thereafter to drink more fluid, though less than controls, whereas alcohol consumption by rats studied by the above authors remained static at the 1 h level. The earlier inhibition by disulfiram of alcohol consumption in our study may be due to the more rapid absorption of the drug solution after intraperitoneal, when compared with that of the drug suspension after oral, administration as reported by the above authors.

Both 3-HK and KA were equally effective, whereas 3-HAA caused a stronger inhibition of alcohol consumption. This is somewhat surprising, as this metabolite did not cause a significant increase in blood acetaldehyde concentration (Fig. 3b), even though it caused the strongest inhibition of ALDH activity after administration (Fig. 2a). It is possible that additional factors are involved in the aversive effects of 3-HAA. Alternatively, it is possible that 3-HAA may also exert an inhibitory effect on ethanol oxidation, which, superimposed on its ALDH inhibition, may have led to the small insignificant elevation of blood acetaldehyde concentration. These possibilities require investigation.

Alcohol consumption was also decreased in alcohol-prefering C57 mice. In particular, the % preference, and to a lesser extent the absolute amount of alcohol consumed, were decreased, with 3-HAA causing the strongest decreases. However, due to the wide individual variations between mice and the small numbers used, these results, though complementary to the aversion results, can only be considered preliminary.

**General safety of kynurenine metabolites**

The present results have shown that a 10 mg/kg body wt dose of the three kynurenine metabolites studied exerted no undesirable side effects or toxicity in rats or mice when administered intraperitoneally once or once daily over a 4–8 day period. KA is the endogenous antagonist of the N-methyl-D-aspartate type of glutamate receptors (Stone, 1993) and doses of it as large as 500–1500 mg/kg body wt are tolerated by mice when administered subcutaneously; only sedation has been observed with these large doses (Rasmussen et al., 1991). Relatively smaller intraperitoneal doses of KA (up to 200 mg/kg) have been shown to be anxiolytic (Lapin, 1996, 1998). The potential toxicity of 3-HK and 3-HAA is a more controversial issue. 3-HAA has been reported (Morita et al., 2001) to induce apoptosis in monocyte-derived cells stimulated by interferon-γ, but at a concentration (200 μM) unlikely to be reached physiologically or even pathologically. Also this effect is limited to the THP-1 and U937, but not four other, cell lines. Under these conditions, the above authors could not detect an apoptotic effect for 3-HK at a similar concentration. In fact, a 250 mg/kg dose of 3-HK has been administered subcutaneously to rats with no reported side effects (Luthman et al., 1996). The fact that our rats treated daily with 3-HK for 8 days gained weight at a rate similar to controls suggests that this metabolite does not cause signs of toxicity at the dose level used. Only 3-HAA caused a significant weight loss, suggesting some degree of toxicity at the 10 mg/kg dose level used, although this was not visibly apparent.

3-HK, which can be taken-up by the brain (Speciale and Schwarz, 1990; Fukui et al., 1991), has been suggested to be neurotoxic, as it can promote neuronal cell death by producing hydrogen peroxide (Okuda et al., 1996) (an effect that has not so far been duplicated *in vivo*) and in view of the earlier finding of its elevated levels in post-mortem brains of patients with Huntington’s disease (Reynolds and Pearsons, 1993), although levels of 3-HK and 3-HAA in blood of living Huntington’s patients (Stoy et al., 2005) and
also those with chronic brain injury (Mackay et al., 2005) have been shown to be lower than in controls. Stay et al. (2005) suggested that, although blood [3-HK] may not reflect that in brain, it is also possible that oxidative stress in Huntington’s disease may trigger a secondary activation of the kynurenine pathway in the brain. Also, although the find-
ings by Okuda et al. (1996) support a pro-oxidant effect of 3-HK, it has been reported (Goda et al., 1999) that, of all Trp and serotonin metabolites, both 3-HK and 3-HAA possess the highest antioxidant properties, with a radical-scavenging reactivity and inhibition of lipid peroxi-
dation greater than those by α-tocopherol (vitamin E). Furthermore, as Morita et al. (2001) suggested that the apop-
totic effect of 3-HAA is due to generation of H$_2$O$_2$, the failure of 3-HK to induce apoptosis in their study further suggests that this kynurenine metabolite does not generate H$_2$O$_2$ even at a 200 μM concentration. Nor are hydroxyl rad-
icals involved in 3-HK oxidation by methaemoglobin with H$_2$O$_2$ (Ishii et al., 1992). Further evidence of an antioxidant role for 3-HK and 3-HAA has been obtained in vitro in rat cerebral cortex and cultured C6 glioma cells with the demon-
stration (Leipnitz et al., 2007) that both metabolites prevent lipid peroxidation in the brain, decrease peroxo radical induction, and prevent glutaric acid-induced free radical for-
mation. Furthermore, the role of 3-HAA as a modulator of the immune system is increasingly recognized with, among others, its ability to decrease the release of cytokines, to exert direct antiproliferative effects, suppress the activation of pro-inflammatory transcription factor NFκB (nuclear factor kappa-light-chain-enhancer of activated B cells), and inhibit nitric oxide synthase (for review, see Darlington et al., 2010 and references cited therein).

General conclusions and comments
We have established proofs of mechanism (inhibition of the target enzyme ALDH after administration and in vivo) and principle (induction of aversion to alcohol) in the present preclinical development study with metabolites of the amino acid Trp of the kynurenine pathway. Previous studies (Lapin et al., 1991) have established a prior art for the use of KA in alcoholism treatment, whereas no prior art exists for the use of 3-HK or 3-HAA, for which a US patent is expected shortly. As the most potent AIDH inhibitor in vivo and the immediate precursor of 3-HAA, 3-HK is of particular interest, but its potential clinical use in alcoholism treat-
ment requires further developmental studies by interested parties. A more immediate and near clinic-ready alternative is the use of its parent compound Trp under appropriate metabolic conditions, evidence for which is presented in the following paper (Badawy et al., 2011).

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