A common East Asian-specific ALDH2 mutation causes obesity and insulin resistance: therapeutic effect of reducing toxic aldehydes by ALDH2 activation

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Abstract

Obesity and type 2 diabetes have reached pandemic proportion. In particular, the population with diabetes is expected to rise rapidly in East and South Asia. ALDH2 (acetaldehyde dehydrogenase 2, mitochondrial) is the key metabolizing enzyme of acetaldehyde and other toxic aldehydes, such as 4-hydroxynonenal (4-HNE).

A missense mutation, Glu504Lys of ALDH2 (denoted as the ALDH2*2 allele) is prevalent in 560 million East Asians, resulting in reduced ALDH2 enzymatic activity. We found that Aldh2*2/*2 homozygous knock-in (KI) mice mimicking human Glu504Lys mutation were prone to develop diet-induced obesity, glucose intolerance, insulin resistance, and fatty liver on a high-fat high-sucrose diet compared with controls. The Aldh2 KI mice demonstrated reduced energy expenditure and thermogenesis. Proteomic analyses of the brown adipose tissue (BAT) of the Aldh2 KI mice identified increased 4-HNE-adducted proteins involved in fatty acid oxidation and electron transport chain. Fatty acid oxidation rate and mitochondrial electron transport activity were reduced in the BAT of the Aldh2 KI mice, which explained the decrease in thermogenesis and energy expenditure.

AD-9308 is a water-soluble prodrug of a potent and highly selective ALDH2 activator AD-5591. In vitro, AD-5591 enhanced both WT and mutant ALDH2 enzymatic activities. AD-9308 allosterically activates ALDH2 mainly by partially blocking the substrate exit tunnel, thereby accelerating the substrate-enzyme collision. In vivo, AD-9308 treatment reduced serum 4-HNE levels, ameliorated diet-induced obesity and fatty liver, and improved glucose homeostasis in both Aldh2 WT and KI mice dose-dependently. Our data highlight the therapeutic potential of reducing toxic aldehyde levels by activating ALDH2 for treating metabolic diseases.

Introduction

Obesity and type 2 diabetes have reached pandemic levels. The 2019 International Diabetes Federation reported that there were 463 million people living with diabetes worldwide. This number is projected to increase rapidly to 578 million in 2030. East Asia and South Asia, the most populous regions in the world, are expected to be the major contributors of new cases. This pandemic is primarily caused by a high-calorie diet enriched in fat and sugar, sedentary lifestyle, and their interaction with genetic factors.

Approximately 36% of East Asians (560 million) or nearly 8% of the global population, carry an inactivating missense mutation of the ALDH2 gene (denoted as the ALDH2*2 allele), which causes Glu504Lys substitution. This mutation results in a reduction of the ALDH2 enzymatic activity by 60–80% in heterozygous carriers (ALDH2*1/*2) and ~90% in homozygous carriers (ALDH2*2/*2). ALDH2*2 heterozygotes and ALDH2*2 homozygotes exhibit sensitivity to alcohol, with presentations ranging from facial flushing, headache, and tachycardia due to a rapid increase in circulating acetaldehyde concentrations. Epidemiological studies suggest a correlation between this inactivating mutation and
several diseases including oral cancer, esophageal cancers, and blood and solid tumors associated with Fanconi anemia. Importantly, several large-scale meta-analyses of genome-wide association studies revealed that this inactivating mutation is strongly associated with type 2 diabetes, body mass index, and serum lipids in East Asians. A validation study further confirmed a close association of ALDH2*2 with visceral fat distribution in 2,958 Chinese subjects. In addition, carriers of the inactivating ALDH2*2 variant had 2–3 times increased risk of non-alcoholic fatty liver disease among Japanese subjects.

The ALDH2 gene encodes mitochondrial aldehyde dehydrogenase 2 (ALDH2), a major acetaldehyde-metabolizing enzyme responsible for ~ 95% acetaldehyde metabolism due to its low Km for acetaldehyde. Acetaldehyde is produced by alcohol dehydrogenase (ADH) following alcohol ingestion. Acetaldehyde can also be generated endogenously from the intermediate metabolisms or by gut microbial flora. Air pollution, thermal degradation of plastics, and spoiled food are also sources of toxic aldehyde. Acetaldehyde is listed as group I carcinogen by the International Agency for Research on Cancer. In addition to acetaldehyde, ALDH2 also metabolizes various bioactive toxic aldehydes, including acrolein, malondialdehyde and 4-hydroxynonenal (4-HNE). Among these bioactive aldehydes, the most intensively studied has been 4-HNE, a lipid peroxidation product that forms covalent adduct to macromolecules such as protein, lipid and DNA, causing cellular damage.

With the advent of high-throughput screening, a small molecule, Alda-1, was identified as an activator for both the wild-type ALDH2 and mutant ALDH2 enzymes. Based on X-ray crystallography and enzyme kinetics, the binding of Alda-1 partially blocks the exit tunnel of substrates, thus increasing the likelihood of productive encounters between reaction intermediate and the catalytic site. Administration of Alda-1 has been effective to treat myocardial infarction, aortic aneurysm, atrial fibillation, Alzheimer’s disease and nociceptive pain in mice model by enhancing the clearance of 4-HNE and in rodent model. The discovery of ALDH2 activators offers an opportunity to test the therapeutic potential of reducing toxic aldehydes for treating a variety of diseases.

AD-9308 is a highly water soluble and orally bioavailable prodrug of a potent and highly selective ALDH2 activator AD-5591, a new generation ALDH2 activator that has the improved biological activities and pharmacological properties compared to Alda-1.

In this study, we used aldh2*2/*2 homozygous knock-in (KI) mice, which mimic the East Asian-specific Glu504Lys mutation, to evaluate the effect of this mutation on diet-induced obesity, glucose homeostasis, fatty liver, and serum lipids and tested the therapeutic effect of a novel ALDH2 activator AD-5591 by dosing its prodrug AD-9308 for treating metabolic disorders.

Results

Aldh2 KI mice carrying the East Asian-specific Glu504Lys mutation were prone to develop diet-induced obesity and fatty liver.
To assess the effect of the East Asian-specific ALDH2 Glu504Lys mutation on diet-induced obesity and related metabolic traits, Aldh2 KI and WT mice littermates were placed on either regular chow or high-fat high-sucrose diet (HFHSD) for 24 weeks since the age of 4 weeks. At the end of study, the body weights of Aldh2 KI and WT mice were 42.45 ± 6.59 and 37.78 ± 5.54 g respectively (Fig. 1a, P < 0.0001). However, no difference of body weight gain was observed between Aldh2 KI and WT mice fed a chow diet (Fig. 1b).

Aldh2 KI mice fed on HFHSD also had more white fat, including inguinal (1.33 ± 0.1 vs. 1.02 ± 0.09 g, P = 0.04), mesenteric (0.68 ± 0.091 vs. 0.51 ± 0.051 g, P = 0.077), and perigonadal fat (2.29 ± 0.27 vs. 1.48 ± 0.13 g, P = 0.004) and increased liver weight (1.89 ± 0.07 vs. 1.64 ± 0.06 g, P < 0.01), but less brown adipose tissue (BAT) weight (0.10 ± 0.01 vs. 0.14 ± 0.10 g, P = 0.016) compared with WT littermates (Fig. 1c). No such differences were found between Aldh2 KI and WT mice on chow diet except for the smaller BAT found in Aldh2 KI mice. Body composition analysis revealed significantly increased fat mass (12.46 ± 0.3 vs. 8.85 ± 1.23 g, P = 0.04), and to a lesser extent, increased lean mass and total water conten in Aldh2 KI mice compared with the WT mice fed on HFHSD (Fig. 1d). Figure 1e showed the representative gross appearance of mice, BAT, perigonadal fat, and liver. H&E stain using of perigonadal fat showed hypertrophic adipocytes with more crown-like necrosis in Aldh2 KI mice compared with controls (Fig. 1f, 1h). There was no difference in the number of adipocytes between the two groups (Fig. 1f-h), indicating hypertrophy rather than hyperplasia of white adipose tissue. Aldh2 KI mice also had significantly higher hepatic triglycerides contents (0.249 ± 0.055 vs. 0.124 ± 0.027 mg/mg liver tissue, P = 0.005), and more severe hepatic steatosis than the WT mice on HFHSD (Fig. 1i, 1j). However, there was no difference in muscle triglycerides content (Fig. 1k). These results suggest that the East Asian-specific ALDH2 Glu504Lys mutation promoted HFHSD-induced obesity and fatty liver in mice.

Aldh2 KI mice had reduced energy expenditure and impaired adaptive thermogenesis

Energy expenditure measured by indirect calorimetry showed that Aldh2 KI mice had lower energy expenditure (Fig. 2a) than WT mice. There was no difference in food intake between Aldh2 KI and WT mice (Fig. 2b). These data indicated Aldh2 KI mice were prone to diet-induced obesity due to reduced energy expenditure but not intake.

Energy expenditure is composed of basal metabolic rate, physical activity, and adaptive thermogenesis including cold-induced and diet-induced thermogenesis. Indirect calorimetry showed no difference in basal metabolic rate as indicated by the energy expenditure in resting (light) phase (Fig. 2a). We also measured energy expenditure by monitoring physical activities of the mice. Daily physical activity, including wheel rotations (Fig. 2c) and travel distances (Fig. 2d) were similar between the two groups. HomeCage monitoring systems did not detect significant differences in various mouse behaviors including awakening, feeding, hanging, rearing up, resting, twisting, and walking except for slightly increased grooming behavior in Aldh2 KI mice (Fig. 2e).

Consequently, we measured adaptive thermogenesis, including cold-induced and diet-induced thermogenesis. For diet-induced thermogenesis, Aldh2 KI mice exhibited significantly lower rectal temperature 30 minutes after HFHSD feeding, indicating impaired diet-induced thermogenesis (Fig. 2f).
For cold tolerance test, *Aldh2* KI mice had lower rectal temperature after 12 hours of prolonged cold exposure at 4 °C (Fig. 2g), indicating cold intolerance. These data suggest that *Aldh2* KI mice may develop obesity due to reduced energy expenditure resulting from impaired adaptive thermogenesis.

**Aldh2 KI mice displayed reduced insulin sensitivity and impaired glucose tolerance**

Since the *ALDH2* Glu504Lys mutation is reported to be associated to type 2 diabetes in genome-wide association studies in East-Asians (3), we further examined glucose homeostasis in mice. On HFHSD, insulin tolerance test (ITT) showed significantly higher blood glucose levels with 76.5% reduction of the inverse area under curve (AUC) of glucose levels in *Aldh2* KI mice, indicating increased insulin resistance (Fig. 2h). Intraperitoneal glucose tolerance (i.p.GTT) showed significantly higher blood glucose levels with AUC of glucose levels increased by 109% in *Aldh2* KI mice (Fig. 2i). Oral glucose tolerance test (OGTT) also showed significantly higher blood glucose levels in *Aldh2* KI mice; AUC of glucose levels during the test were increased by 111% in KI mice, indicating worsened glucose intolerance (Fig. 2j). *Aldh2* KI mice displayed a compensatory increase in insulin secretion after oral glucose load (Fig. 2k). No differences in glucose homeostasis, measured by glucose and insulin tolerance tests, were found between *Aldh2* KI and WT mice fed a chow diet (Supplementary Fig. 1).

In addition, *Aldh2* KI mice on the HFHSD showed a trend for increased total cholesterol (204.5 ± 7.4 vs. 186.4 ± 7.5 mg/dl, P = 0.09, Fig. 2l), triglycerides levels (107 ± 7.63 vs. 91.33 ± 2.75 mg/dl, P = 0.10) (Fig. 2m) and serum leptin (19.6 ± 2.8 vs. 12.7 ± 1.8 mg/ml, P = 0.15, Fig. 2n). However, adiponectin levels were not different between the two groups (Fig. 2o).

**Aldh2 KI mice had reduced mitochondrial fatty acid oxidation rate and lower respiratory transport chain activity in brown adipose tissue due to 4-HNE adduction**

In view of the reduced adaptive thermogenesis observed in the *Aldh2* KI mice, we compared the expression levels of *Ucp1*, the major thermogenic protein in BAT and white fat including inguinal and perigonadal fat between the *Aldh2* KI and WT mice by real-time quantitative PCR (RT-qPCR). There was no difference in *Ucp1* expression in the BAT or white fat between the *Aldh2* KI and WT mice fed on HFHSD (Fig. 3a). Immunoblots further confirmed that there was no difference in the level of Ucp1 in the BAT, the tissue where Ucp1 was normally expressed (Supplementary Fig. 2).

We next examined the expression of mitochondrial respiratory complex I to V component in BAT. These protein components are essential for the maintenance of the proton gradient in mitochondrial electron transport chain (ETC) and are required for UCP1-mediated thermogenesis. There was no difference in mitochondrial respiratory complex I to V protein components expression in the BAT of *Aldh2* KI and WT mice (Supplementary Fig. 3). We also observed no difference in microscopic morphology of BAT, the major thermogenic organ (Supplementary Fig. 4).

Since the ETC protein expression studied showed no difference between *Aldh2* KI and WT mice, it implied that the decreased energy expenditure and thermogenesis were likely due to impaired protein functions.
We found increased protein carbonylation (Fig. 3b) and 4-HNE-adducted mitochondrial proteins (Fig. 3c) in the BAT of the KI mice compared with WT mice. Using liquid chromatography tandem mass spectrometry (LC MS/MS) analysis, we identified 20 4-HNE-adducted BAT mitochondrial proteins in Aldh2 KI mice and 10 4-HNE adducted mitochondrial proteins in WT mice, with 8 proteins which are present in both Aldh2 KI and WT mice (Fig. 3d, 3e).

Three identified adducted sites were present in proteins involved in fatty acid oxidation (FAO) including 3-ketoacyl-CoA thiolase and propionyl-CoA carboxylase. Eleven sites were present in proteins involved in the maintenance of mitochondrial electron transport chain (ETC) including NADPH dehydrogenase (complex I), succinate dehydrogenase (complex II), cyochrome b-1 complex (complex III), ATP synthase, and glycerol-3-phosphohydrogenase. Three adducted sites were present in the MICOS complex, which are essential for the maintenance of inner mitochondrial membrane structure. Hence, we further evaluated the capacity of mitochondrial FAO and ETC to determine the effect of 4-HNE modification of these proteins. The ex vivo FAO rate of the whole BAT tissue isolated from Aldh2 KI mice was significantly reduced by 16% (P = 0.03) compared with WT mice (Fig. 3f). Consistent with the whole BAT tissue, the FAO rate of the cultured primary brown adipocytes isolated from the Aldh2 KI mice was also significantly reduced by 22% (P = 0.03, Fig. 3g). Furthermore, addition of 4-HNE decreased FAO rate of induced primary brown adipocytes isolated from WT mice in a dose-dependent manner; FAO rates were reduced by 36% (P < 0.01) and 60% at 5 and 10 µM 4-HNE, respectively (P < 0.01) (Fig. 3h). In line with the LC-MS/MS finding, we found a significant reduction in the enzymatic activity of mitochondrial respiratory complexes I, II, and III, but not complex IV, in Aldh2 KI mice compared with WT mice (Fig. 3i). Oxygen consumption rate was also significantly decreased in induced primary brown adipocytes isolated from Aldh2 KI mice compared with WT mice (Fig. 3j). Since FAO is the major energy source of thermogenesis and the mitochondrial ETC is essential for maintaining proton gradient required for thermogenesis, the increased protein carboxylations may explain the impaired thermogenesis of Aldh2 KI mice.

**ALDH2 activator AD-9308/AD-5591 activated wild-type and mutant human ALDH2 enzymatic activity**

AD-9308 is a valine ester prodrug of a potent and selective small molecule ALDH2 activator AD-5591. When administered in vivo, AD-9308 is rapidly converted to AD-5591 by esterase hydrolysis (Fig. 4a). In vitro, AD-5591 treatment increases the catalytic activity of recombinant wild-type and mutant human ALDH2 (Fig. 4b). In vivo, AD-9308 administration showed a good pharmacokinetic profile in mice when administered orally or intravenously with good bioavailability in rodents and dogs (Supplementary Table 2).

Alda-1 allosterically activates ALDH2 mainly by partially blocking the substrate exit tunnel, thereby accelerating the substrate-enzyme collision without impeding the catalytic sites of Cys302 and Glu268; Alda-1 also inhibit substrate-induced ALDH2 inactivation by protecting Cys301 and Cys303 oxidation. Using molecular docking, we found that the binding pocket for AD-5591 (Fig. 4c) is close to that of
Alda-1 (Supplementary Fig. 5a). Similar to Alda-1 (Supplementary Fig. 5b), AD-5591 is bound within a hydrophobic collar by Tyr456, Val458, Lys127, Met124, Gln462, Gly460, Phe459, Val120, and Phe292 of human ALDH2 (Fig. 4d). This binding site leaves the catalytic sites Cys302 and Glu268 unimpeded (Fig. 4e, Supplementary Fig. 5c). Therefore, the mechanism by which AD-5591 activates ALDH2 is very similar to Alda-1. Furthermore, Alda-1 forms a single hydrogen bond with the Asp457 residue of ALDH2. Yet, in our modeling analyses, AD-5591 forms one additional hydrogen bond with Ala461 (Fig. 4e), which may explain its higher affinity to ALDH2 than for Alda-1.

ALDH2 activator AD-9308/AD-5591 lowered 4-HNE and attenuated diet-induced obesity, fatty liver, insulin resistance, and glucose intolerance in both WT and Aldh2 KI mice

In view of the marked reduction in Aldh2 expression in diet-induced obese mice (Supplementary Fig. 6), we next explored whether Aldh2 activation can rescue these obesity-associated phenotypes. From the age of 10 weeks when HFHSD was started, Aldh2 KI and WT mice were treated with vehicle, 20 mg/kg/day or 60 mg/kg/day of AD-9308 by oral gavage for 20 weeks (Fig. 5a). As shown in Fig. 5B and 5C, AD-9308 treatment effectively reduced the diet-induced weight gain in both Aldh2 KI and WT mice dose-dependently. Weight of perigonadal fat, inguinal fat, omental fat, and liver decreased with AD-9308 treatment dose-dependently (Fig. 5d-5 h). AD-9308 treatment reduced the extent of hepatic steatosis (Fig. 5i, 5j) and the levels of hepatic triglycerides contents in both Aldh2 KI and WT mice in a dose-dependent manner (Fig. 5k).

Fasting glucose levels were lowered by AD-9308 treatment in both Aldh2 WT and KI mice (Fig. 6a, 6b). AD-9308 treatment effectively reduced insulin resistance in both WT (Fig. 6c) and KI (Fig. 6d) mice and significantly improved glucose tolerance of both WT (Fig. 6e) and KI (Fig. 6f) mice in a dose-dependent manner. AD-9308 also decreased serum 4-HNE levels in both mice groups dose-dependently (Fig. 6g). Figure 6h. shows the summary diagram depicting how reducing 4-HNE by ALDH2 activator AD-9308 ameliorates metabolic disturbances.

Pathological examination reveled no abnormalities in liver or kidney in both Aldh2 WT and KI mice treated with AD-9308 for 20 weeks (Supplementary Table 3). Serum alanine aminotransferase (ALT) and creatinine levels were also not different between groups after AD-9308 treatment for 20 weeks (Supplementary Fig. 7).

Discussion

The prevalence of obesity and diabetes mellitus has surged in the past decades and is predicted to continue to rise, especially in East and South Asia. This trend is largely caused by high-calorie diet enriched in fat and sugar, sedentary lifestyle, and their interaction with genetic predisposition. Genome-wide association studies have confirmed many genetic loci associated type 2 diabetes and obesity. Specifically, several genetic loci are East Asian-specific. Genetic variants in or near the CDKAL1, KLF9, GP2, ALDH2, and ITIH4 genes are associated with obesity and genetic variants in or near the
GDAP1, PTF1A, SIX3, ALDH2, and PAX4 genes\textsuperscript{3,34} are specifically associated with type 2 diabetes in East Asians. Among them, the ALDH2 Glu504Lys mutation that affects 560 million East Asians or nearly 8% of global population is associated with body mass index, type 2 diabetes, and serum lipids in large meta-analyses of genome-wide association studies.

We demonstrated that Aldh2 knock-in mice carrying the East Asian-specific Glu504Lys mutation were more prone to develop diet-induced obesity, fatty liver, insulin resistance and glucose intolerance than WT mice on HFHSD. Importantly, the ALDH2 activator AD-9308 increased both the catalytic activity of WT and mutant enzyme, reduce serum 4-HNE levels, and effectively alleviated diet-induced obesity, fatty liver, insulin resistance, and glucose intolerance in both Aldh2 KI and WT mice in a dose-dependent manner.

BAT is a highly specialized organ enriched in Ucp1 for adaptive thermogenesis. Although Aldh2 KI mice exhibited impaired thermogenesis, unexpectedly, they did not show reduced Ucp1 expression. Instead, we found that several key mitochondrial proteins involved in mitochondrial FAO and ETC were modified by 4-HNE adduction. Consistently, previous studies have shown that 4-HNE is mainly generated from oxidation of mitochondrial membranes, with 30% of 4-HNE-adducted proteins located within mitochondria\textsuperscript{35,36}.

Our data further support that the 4-HNE adduction to mitochondrial proteins can be an underlying mechanism of ALDH2 inactivation that leads to the reduction of mitochondrial FAO and ETC function.

Mitochondrial FAO and ETC are required for the maintenance of the proton gradient in the intermembranous space, which is essential for Ucp1-mediated adaptive thermogenesis. In our study, we found that the thermogenic capacity of Aldh2 KI mice was reduced. Fatty acids serve as the main fuel suppliers for thermogenesis\textsuperscript{37}. It has been estimated that intracellular fatty acids in the BAT contribute 74–84% of the fuel for thermogenesis upon cold challenge\textsuperscript{37}. Cpt1 is the rate-limiting enzyme for the translocation of fatty acids into mitochondria for \(\beta\)-oxidation. Cpt1b\textsuperscript{+/-} mice developed fatal hypothermia following cold challenge\textsuperscript{38}. Adipose-specific Cpt2-knockout mice presented a hypothermic phenotype when exposed to cold\textsuperscript{39}. Mice deficient in fatty acid \(\beta\)-oxidation enzymes, including very-long-chain acyl-CoA dehydrogenase (VLCAD), long-chain acyl CoA dehydrogenase (LCAD), and short-chain acyl CoA dehydrogenase (SCAD) also displayed cold intolerance\textsuperscript{40–42}. These data indicate that mitochondrial FAO is critical for adaptive thermogenesis. Furthermore, BAT-specific Lkb1-knockout mice, which have reduced expression of ETC complex proteins, also developed impaired thermogenesis\textsuperscript{43}, indicating that the integrity of mitochondrial ETC machinery is essential for adaptive thermogenesis. These data strongly support our findings that 4-HNE adduction to mitochondrial proteins involved in mitochondrial FAO and ETC could lead to impaired adaptive thermogenesis.

ALDH2 metabolizes bioactive toxic aldehydes by oxidation. In addition to ALDH2, one of the major pathways to detoxify 4-HNE is mediated through glutathione transferases (GSTs) by conjugation to glutathione. Gsta4 is one of the isoforms of GST with the highest conjugation activity for 4-HNE. Consistent with the present finding, disruption of Gsta4 in mice increased 4-HNE levels and caused obesity in mice\textsuperscript{44}. Disruption of the Gst-10 gene, which causes a 50% increase in 4-HNE adducts in C.
*elegans* also resulted in fat accumulation and direct treatment with 4-HNE increases lipid storage in *C. elegans*. Conversely, over-expression of *Gst-10* led to 4-HNE reduction and a lean phenotype. Glutathione peroxidase 4 (*Gpx4*), which resides in the inner mitochondrial membrane, is a scavenger that reduces lipid peroxides. Deficiency of *Gpx4* in mice increased the number of 4-HNE adducts and exacerbated glucose intolerance, dyslipidemia, and fatty liver. Collectively, these data indicate that increased aldehydic load due to excessive 4-HNE cause metabolic disorders. Indeed, overexpression of *ALDH2* has been shown to decrease both the heart and liver weight and mitochondrial injury in mice fed on a high-fat diet. Interestingly, diet-induced obesity was not observed in these *ALDH2* over-expressing mice.

Our study also showed that the *Aldh2* KI mice had more severe hepatic steatosis than the WT mice when fed on HFHSD, which can be reversed by AD-9308 treatment. In line with the present results, Alda-1, a prototype of the ALDH2 activator has been shown to alleviate nonalcoholic hepatic steatosis in apolipoprotein E-knockout mice and reversed alcohol-induced hepatic steatosis in animals, supporting activation of ALDH2 also prevent both alcohol and non-alcoholic hepatic steatosis.

Our study has several limitations. First, obesity and related metabolic phenotypes in the *Aldh2* KI mice were observed only when fed on HFHSD and not when placed on chow diet, the metabolic phenotypes were not different between the *Aldh2* KI and WT mice. Therefore, our findings may not be generalized to the entire population. The HFHSD for mice consists of 58% calories from fat and 12% calories from sucrose; while the chow diet is composed of 13% calories from fat and 3% calorie from sucrose. According to the Nutrition and Health Surveys in Taiwan (NAHSIT) in 2008, the average calorie intake from fat and sucrose is 33% and 8% in adults in Taiwan. In the National Health and Nutrition Examination Survey of U.S adults., the average calorie intake from fat and sucrose is 35% and 14% in 2012. The sucrose content is comparable between HFHSD for mice and modern human diet but the fat content of HFHSD is higher than human diet. Even if this concern is relevant, the findings of this study are still substantial given the large number of East Asians (560 million people) carrying this inactivating mutation (Glu504Lys) in the *ALDH2* gene. Second, our experiments did not include heterozygous knock-in mice. Third, although the metabolic disorders were normalized in mice treated with AD-9308 for 5 months without pathological changes in the liver and kidney, long-term safety should be formally determined with GLP standard. Last, humans and other mammals have 19 different aldehyde dehydrogenases (ALDH) and at least six are found in the mitochondria. Although AD-9308 does not activate ALDH1A1, ALDH3A1, ALDH4A1, ALDH5A1 and ALDH7A1 members of the ALDH family (data not shown), it is possible that AD-9308 may still have non-specific effect on other ALDHs.

Our study has important clinical implications. We showed that reduced activity of the mitochondrial enzyme, ALDH2, exacerbates obesity-associated pathologies. These pathologies correlate with increased aldehydic load and inactivation of critical mitochondrial proteins involved in FAO and ETC. Significantly, we showed that treatment with an activator of ALDH2, such as AD-9308 prevented these pathologies. These data provide strong evidence for a critical role of toxic aldehydes accumulation and defective...
ALDH2 activity in the pathogenesis of obesity, diabetes, and fatty liver disease. Our study provide a new strategy by targeting ALDH2 for the treatment of obesity-associated metabolic disorders which is rising rapidly in human populations, particularly in the East and South Asia.

Methods

Animal experiments, administration of diets and drug treatment

All animal experiments were performed according to institutional ethical guidelines and were approved by the Institutional Animal Care and Use Committee (IACUC), which is accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International (AAALAC). The mice were housed at 22–24 °C with light: dark cycles of 12:12 hours. The mice were fed either a HFHSD (cat. no. D12331, Research Diets) which provided 58% calorie from fat and 12.5% calorie from sucrose or chow diet (cat no. 5001, Lab Diet). For the animal experiments, AD-9308 was dissolved in water and was delivered daily to mice by oral gavage. Only male mice were used in this study.

Generation of Aldh2*2/*2 knock-in mice

Aldh2 KI mice carrying the human ALDH2 Glu504Lys mutation were generated by introducing the Glu504Lys mutation within the mouse gene. Both Aldh2 WT controls and Aldh2 KI mice were littermates from mated heterozygous mice.

Glucose and insulin tolerance test

Oral and intraperitoneal GTT were evaluated after 6 hours of fasting. Tail blood glucose was collected at 0, 15, 30, 45, 60, 90, and 120 min after oral gavage or intraperitoneal injection of glucose water (1 g/kg) and measured by a glucometer (ACCU-CHECK Performa, Roche). For the ITT, mice were intraperitoneally injected 1 U/kg insulin (Humulin R, Eli Lilly) after 4 hours of fasting. Tail blood glucose was collected at 0, 15, 30, 45, 60, 90, 120, and 180 min after injection.

Energy expenditure, food intake and physical activity

Metabolic measurements (food and water intake, locomotor activity, VO_{2} consumption and VCO_{2} production) were obtained using the Promethion metabolic phenotyping system (Sable Systems). Monitoring was performed for 5 days after mice have been acclimatized to the cages for 2 days.

Hepatic triglycerides content measurement

Approximately 80 mg of liver tissue was homogenized in 1800 µl of chloroform/methanol (2:1). Then, 360 µl of H_{2}O was added. The homogenates were centrifuged and the lower 200-µl layer was added with 100 µl of chloroform with 4% Triton X-100 and then dried in a chemical hood. The dried pellet was redissolved with 200 µl of H_{2}O for determination of triglyceride concentrations with Wako TG LabAssay kit (cat. no. 290-63701, Wako)
Cold tolerance test and diet-induced thermogenesis test

For the cold tolerance test, 24-week-old mice with matched average body weight from the two groups were placed individually on HFHSD in a 4°C chamber. The rectal temperature of the mice was measured after 0, 1, 2, 3, 4, 5, 6, 12, and 18 hours. For the diet-induced thermogenesis test, 24-week-old mice were fasted overnight for 18 hours. Then, their rectal temperature was measured at 0, 15, 30, 60, 120, 180, and 240 min after HFHSD refeeding.

Real-time quantitative PCR (RT-qPCR)

RT-qPCR was performed using SYBR green reagent (cat. no. 11203ES08, YEASEN). The primer sequences for mouse Ucp1 and Ppia is list in Table S1. RT-qPCR reactions were performed using an ABI 7900HT FAST (Applied Biosystems). All qPCR reactions were run in duplicate for each sample.

Primary brown adipocyte culture

BAT from Aldh2 WT and KI mice aged 4 weeks was minced and digested by type I collagenase (cat. no. 17018029, Thermo Fisher Scientific) in HEPES buffer. The stromal vascular fraction (SVF) was obtained by centrifugation and cultured in Dulbecco’s modified Eagle medium:nutrient mixture F-12 (DMEM/F-12) (cat. no. 12500062, HyClone) supplemented with 10% FBS (cat. no. 04-001-1A, Biological Industries) and 1% antibiotic/antimycotic solution (cat. no. SV30079.01, HyClone). For cell differentiation, preadipocytes were cultured in differentiation medium containing 10% FBS, 0.5 mM isobutyl-methylxanthine, 1 µg/ml insulin, 5 µM dexamethasone, 1 nM T3, and 125 µM indomethacin for 2 days. Next, cells were maintained in the medium containing DMEM/F12 with 10% FBS, 1 µg/ml insulin, and 1 nM T3. The medium was changed every 2 days.

Isolation of mitochondria from brown adipose tissue

BAT was removed from Aldh2 IWT and KI mice fed a HFHSD for 5 weeks. Isolation of mitochondria was performed according to published protocols 53. Briefly, BAT was homogenized in 10% w/v of ice-cold mitochondrial isolation buffer by a Dounce homogenizer. The pellets were separated by centrifugation, and then resuspended in SET buffer (0.25 M sucrose, 1 mM EDTA, and 10 mM Tris-HCl, pH 7.4) for the fatty acid oxidation assay and LC-MS/MS analysis or in BES-sucrose buffer (0.25 M sucrose, 0.1 M KCl, and 20 mM BSE, pH 7.2) for the mitochondrial complex activity assay.

Fatty acid oxidation (FAO) assay

FAO measurements were performed using labeled 3H-palmitic acid and 3H2O production was assessed as previously described 54. For FAO rate of differentiated primary cells, the capture of 3H2O was measured after a 2-hour incubation with 5 mM palmitate/BSA buffer including 0.5 µCi [3H]-palmitate (cat. no. PK-NET043001MC, PerkinElmer) in the presence of 1 mM carnitine. For FAO rate of whole BAT, the isolated mitochondria were placed in a 24-well plate and added incubated with reaction buffer (100 mM sucrose, 10 mM Tris-HCl, 5 mM KH2PO4, 0.2 mM EDTA, 80 mM KCl, 1 mM MgCl2, 2 mM L-carnitine, 0.1 mM malate, 0.05 mM coenzyme A, 2 mM ATP, 1 mM dithiothreitol, and 7% BSA/5 mM palmitate/0.01 µCi/µl
[3H]-palmitate) at 37°C for 60 min. 3H2O was isolated by oil-water separation with chloroform, methanol and KCl/HCl. The average counts per minute (CPM) were measured using a liquid scintillation counter.

**Biotin hydrazide staining for carbonylated protein**

For detection of carbonylated protein, samples were chemically reduced by NaBH4 and then incubated with 0.5 mM EZ-link biotin hydrazide (ca. no.21339, Pierce) for 1 hour. After coupling, the samples were separated by 10% SDS-PAGE gel and transferred to PVDF membrane. The membrane was blocked with 10% skim milk in PBS containing 0.05% Tween-20 (PBST) at 4°C overnight and then incubated with Streptavidin-HRP (cat no. 890803, BD Biosciences) for 1 hour at room temperature. Chemiluminescence signals were developed with HRP substrate (cat. no.WBLUR0500, Millipore).

**Plasmid construction, expression and purification of human ALDH2 in Escherichia coli**

*E. coli* BL21 (DE3) was transformed with pTrcHi-WT ALDH2 and pTrcHi -KI ALDH2. The transformants were cultured and then induced to express recombinant proteins using 0.4 mM IPTG for 16–18 hours at 25 °C. The cells were harvested by centrifugation and broken in lysis buffer by sonication on ice. The recombinant ALDH2 protein was purified with Ni-NTA resin (cat. no. 88222, ThermoFisher) following the user manual.

**ALDH2 enzymatic activity assays**

ALDH2 activity was measured by monitoring the reduction rate of NAD+/min at 340 nm and 25 °C. Enzyme activity was assayed in 100 µl of reaction mixtures containing 50 mM Na4P2O7 (pH 9.5), 0.01% BSA, 10 mM NAD+, 50 µM acetaldehyde and recombinant ALDH2 with or without 100 µM AD-5591 or Alda-1. One enzymatic activity unit was defined as the reduction of 1 µmol NAD+ per min by 1 mg of human ALDH2 protein.

**Molecular docking**

To visualize the interaction between the ALDH2 with activators Alda-1 and AD-5591, the X-ray structure of human ALDH2 (PDB ID: 1NZX) and NAD+ were used. Ligand energy was minimized using the PyRx program before docking. Three-dimensional models are generated by using the PyMOL program and the 2D protein-ligand interaction diagrams were generated by the LigPlot + program.

**LC – MS/MS analysis of 4-HNE-modified proteins**

The protein samples were prepared with in-solution trypsin digestion. The peptides were desalted with C18 Zip Tip (Millipore, USA) and dry by vacuum centrifugation. The desalted and dried peptides were resuspended in 0.1% formic acid.

LC-MS/MS analysis was performed on a NanoACQUITY UPLC system (Waters, USA) coupled to a high-resolution mass spectrometer (QE HF-X, Thermo Fisher Scientific). The peptides were injected into a trap column (Symmetry C18, 2 cm × 75 µm i.d.) and then separated in a 25 cm × 75 µm i.d. BEH130 C18
column (Waters, USA) on a gradient from 0–85% buffer B (buffer A, 0.1% FA H₂O; buffer B, 0.1% formic acid in acetonitrile). The mass spectrometer was operated in data-dependent mode with the following acquisition cycle: an MS scan (m/z 350–1600) recorded at resolution R = 60,000 and MS/MS scans recorded at resolution R = 15,000, which were acquired by HCD fragmentation with collision energy of 28.

MS/MS spectra were searched with the Mascot engine (v2.6, Matrix Science) against the UniProtKB mouse protein database using the following parameters: a precursor peptide mass tolerance of 20 ppm and an MS/MS fragment tolerance of 0.02 Da with a maximum of two missed cleavage sites. The following modifications were made to the peptides: static carbamidomethylation on cysteine, variable oxidation on methionine, variable deamidation of asparagine or glutamine, and various 4-HNE modifications on cysteine, histidine and lysine. The cut-off threshold for acquiring significant peptide-to-spectrum matches was P < 0.05.

**Mitochondrial respiratory chain complex activity assay**

The mitochondrial complex spectrophotometric assays were carried out using published protocols. Complex I and complex II activities were measured spectrophotometrically by examining the decrease in absorbance due to the reduction of 2,6-dichlorophenolindophenol (DCPIP) at 600 nm. Activity was expressed in nanomoles of DCPIP reduction/min/mg protein (E = 19.1 mmol⁻¹·cm⁻¹). Complex III-specific activity and complex IV-specific activity were measured by monitoring the reduction of oxidized cytochrome C(III) and oxidation of reduced cytochrome C(II) at 550 nm, respectively. The activity is expressed as a nanomole of reduced cytochrome of oxidized cytochrome C /min/mg protein (E = 18.5 mmol⁻¹·cm⁻¹).

**Immunoblots to detect 4-HNE-adducted proteins**

Samples prepared with Laemmli sample buffer and were separated by 10% SDS-PAGE gel and transferred to PVDF membrane. The membrane was blocked with 10% skim milk in PBST and incubated at 4°C overnight with anti-4-HNE antibody (1:1000; cat. no. PAB1295, Abnova) and then with secondary antibodies with HRP (1:10000; cat. no. GTX26721, GeneTex).

**Oxygen consumption rate (OCR)**

Stromal vascular fraction was isolated from BAT of 4-week-old Aldh2 WT and KI mice and seeded to Seahorse XF24 v7 cell culture plates (Agilent) at approximate density of 20,000 cells per well and was then differentiated to primary brown adipocyte as described above. On day7 of differentiation, cells were washed once with Seahorse medium and maintained in Seahorse medium in CO₂-less incubators. Basal OCR were determined by three measurements, followed by three measurements after each injection drug: oligomycin (2 µM), FCCP (2 µM), Rotenone/Antimycin A (0.5 µM) using Seahorse XFe24 Analyzer (Agilent). OCR measurements of each stage were normalized to the protein concentration.

**Serum biochemistry**
Blood was collected from mice fasted for 4 hours. Fasting plasma adiponectin (cat. no. ab108785, Abcam), leptin (cat. no. DY498, R&D), and insulin (cat. no. 10-1247-01, Mercodia), 4-HNE (cat. no. EEL-M2677, Elabcience) concentrations were measured using ELISA kits. Plasma triglycerides, and total cholesterol levels were measured using the FUJI DRI-CHEM clinical chemistry analyzer.

**Statistical analyses**

Two-tailed independent t-tests were used for comparing two independent groups. The Wilcoxon rank-sum test was used to compare data from RT-qPCR. Statistical analyses were conducted using GraphPad Prism 5.0. All P-values < 0.05 were considered statistically significant.

**Data availability**

All data associated with this study are available in the main text or the supplementary materials

**Declarations**

**Competing interests**


**Author contributions**

H.L.L. performed the LC-MS/MS analyses, immunoblots, RT-qPCR, histology examinations, and enzyme activity assays. J.Y.H. and W.L.S. performed the animal experiments. Y.C.C and J.Y.N. performed ELISA. Z.Z.D. performed histology examinations. M.L.H. and Y.T.T. conducted the pathological examinations. M.L.H. performed molecular docking. C.C.L. performed the β-oxidation assay and LC-MS/MS experiments. T.Y.L. performed the OCR measurement. W.Y designed and synthesized AD9308 and AD5591 and conducted pharmacokinetic studies. F.A.L. performed the LC-MS/MS experiments. H.L.C. performed indirect calorimetry. L.Y.C. drew summary graphs. Y.C.C. H.L.L, W.Y, C.H.C. D.M.R. and L.M.C. wrote the manuscript. L.M.C. D.M.R. and Y.C.C. conceived the study and participated in the design and interpretation of all experiments.

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References


Figures
Figure 1

Body weight of Aldh2 WT and KI mice on (a) HFHSD (n=24:21) and (b) chow (n=15:19). (c) Weights of inguinal, perigonadal, and mesenteric fat, the liver, and brown adipose tissue (BAT) of the Aldh2 WT and KI mice on HFHSD (n=67:48) and chow (n=10:12). (d) Body composition of Aldh2 WT and KI mice on a HFHSD (n=12:20) and chow (n=15:19). (e) Gross appearance if mice, BAT, perigonadal fat and liver. (f) Average adipocyte size and (g) adipocyte number in the perigonadal fat pad of the Aldh2 WT and KI mice on HFHSD (n=24:16). (h) H&E stain of the perigonadal fat in Aldh2 WT and KI mice on HFHSD. (i) Hepatic triglyceride content of the Aldh2 WT and KI mice on HFHSD (n=23:20) and chow diet (n=10:8). (j) Representative H&E staining of livers from the Aldh2 WT and KI mice. (k) Muscle triglyceride content in the Aldh2 WT and KI mice on HFHSD (n=23:20) and chow diet (n=10:8). *P < 0.05, **P< 0.01.
Figure 2

(a) Energy expenditure (n=12:12), (b) food intake (n=12:12), (c) wheel rotations (n=12:12), (d) distance traveled (n=6:6), and (e) behaviors monitored by HomeCage systems (n=6:6) of the Aldh2 WT and KI mice on HFHSD. (f) Diet-induced thermogenesis and (g) cold-induced thermogenesis of the Aldh2 WT and KI mice on a HFHSD (n=28:18). (h) Glycemic levels during the insulin sensitivity test (n=50:44) and (i) intraperitoneal glucose tolerance test (n=50:44), and (j) glycemic levels during the oral glucose tolerance
test of Aldh2 WT and KI mice on HFHSD (n=50:44). (k) Insulin levels following oral glucose load (n=45:36), and fasting serum (l) total cholesterol, (m) triglycerides, (n) leptin, and (o) adiponectin (n=24:21) of Aldh2 WT and KI mice on HFHSD. *P < 0.05, **P< 0.01.

**Figure 3**

(a) Ucp1 expression levels in inguinal fat, perigonadal fat, and brown adipose tissue (BAT) of the Aldh2 WT and KI mice measured by RT-qPCR (n=21:17). (b) Protein carbonylation of BAT mitochondria by
hydrazide biotin staining. (c) Immunoblots of 4-hydroxynonenal (4-HNE)-adducted mitochondrial proteins in the BAT from the Aldh2 WT and KI mice on HFHSD. (d,e) 4-HNE-adducted mitochondrial proteins and amino acid residues of the BAT from the Aldh2 KI and WT mice, as identified by LC-MS/MS (n=2:2). (f) Fatty acid oxidation (FAO) rate of the whole BAT tissue isolated from the Aldh2 WT and KI mice fed a HFHSD (n=24:20). (g) FAO of primary brown adipocytes isolated from the Aldh2 WT and KI mice fed a HFHSD (n=3 per group). (h) FAO rate of induced primary brown adipocytes isolated from the WT mice treated with 0, 5, and 10 μM 4-HNE (n=3 per group). (i) Mitochondrial electron transfer chain (ETC) complex I-IV enzymatic activity isolated from the BAT of Aldh2 WT and KI mice on HFHSD (n=4:4) (j) Oxygen consumption rate (OCR) of induced primary brown adipocytes isolated from Aldh2 WT and KI mice fed on HFHSD. Cells were treated with oligomycin, FCCP, and Rotenone/Antimycin A. OCR measurements were normalized to the protein concentration. (n=3 per group) *P<0.05, **P< 0.01.
Figure 4
(a) Prodrug AD-9308 is converted to AD-5591 by esterase hydrolysis in vivo. (b) AD-5591 significantly increases the enzymatic activity of recombinant WT and mutant human ALDH2 proteins (n=9:9). (c) Ribbon diagram showing the binding pocket of AD-5591 within human WT ALDH2. (d) LigPlot showing the bonds between AD-5591 and human WT ALDH2. (e) Regional ribbon diagram showing the binding of AD-5591 and NAD+ with WT ALDH2.
Figure 5

(a) Study flow of AD-9308 treatment. Body weight of (b) Aldh2 WT and (c) KI mice treated with vehicle, low-dose AD-9308 (20 mg/kg/day), and high-dose AD-9308 (60 mg/kg/day). (d) Gross appearance of the perigonadal fat from the Aldh2 WT and KI mice treated with AD-9308 on HFHSD. Weight of (e) perigonadal fat, (f) inguinal fat, (g) omental fat, (h) liver of the Aldh2 WT and KI mice treated with AD-9308. (i) Gross appearance of the liver. (j) H&E stained liver, and (k) hepatic triglycerides content of the Aldh2 WT and KI mice treated with AD-9308 on HFHSD (n=11:14 for the vehicle group; n=12:15 for 20 mg/kg/day group; n=9:12 for 60 mg/kg/day group). *P < 0.05 **P < 0.01.
Figure 6

Fasting glucose level of the (a) Aldh2 WT and (b) KI mice treated with vehicle, low-dose AD-9308 (20 mg/kg/day), and high-dose AD-9308 (60 mg/kg/day) on HFHSD. (c)(d) Glycemic levels during the insulin sensitivity test and (e)(f) intraperitoneal glucose tolerance test of the Aldh2 WT and KI mice treated with AD-9308 (n=15:15 for the vehicle group; n=14:16 for 20 mg/kg/day group; n=9:14 for 60 mg/kg/day group). (g) Serum 4-HNE levels of the Aldh2 WT and KI mice treated with AD-9308 (n=15:16 for the vehicle group; n=12:14 for 20 mg/kg/day group; n=12:14 for 60 mg/kg/day group) on HFHSD. (h) Summary diagram depicting how reducing 4-HNE by AD-9308 ameliorates metabolic disturbances. *P < 0.05 **P < 0.01.

Supplementary Files

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