The Mutation in the Mitochondrial Aldehyde Dehydrogenase (ALDH2) Gene Responsible for Alcohol-induced Flushing Increases Turnover of the Enzyme Tetramers in a Dominant Fashion

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Abstract

Deficiency in mitochondrial aldehyde dehydrogenase (ALDH2), a tetrameric enzyme, results from inheriting one or two ALDH2*2 alleles. This allele encodes a protein subunit with a lysine for glutamate substitution at position 487 and is dominant over the wild-type allele, ALDH2*1. The ALDH2*2-encoded subunit (ALDH2K) reduces the activity of ALDH2 enzyme in cell lines expressing the wild-type subunit (ALDH2E). In addition to this effect on the enzyme activity, we now report that ALDH2*2 heterozygotes had lower levels of ALDH2 immunoreactive protein in autopsy liver samples. The half-lives of ALDH2 protein in HeLa cell lines expressing ALDH2*1, ALDH2*2, or both were determined by the rate of loss of immunoreactive protein after inhibition of protein synthesis with puromycin and by pulse-chase experiments. By either measure, ALDH2E enzyme was very stable, with a half-life of at least 22 h. ALDH2K enzyme had an enzyme half-life of only 14 h. In cells expressing both subunits, most of the subunits assemble as heterotetramers, and these enzymes had a half-life of 13 h. Thus, the effect of ALDH2K on enzyme turnover is dominant. These studies indicate that the ALDH2*2 allele exerts its dominant effect both by interfering with the catalytic activity of the enzyme and by increasing its turnover. This represents the first example of a dominantly acting allele with this effect on a mitochondrial enzyme’s turnover. (J. Clin. Invest. 1996; 98:2027–2032.) Key words: aldehyde • ethanol • enzyme • liver • protein degradation

Introduction

The well-known alcohol-induced flush reaction is caused by a mutation in the structural gene for the mitochondrial aldehyde dehydrogenase (ALDH2) (1, 2). The variant allele, designated ALDH2*2, encodes a lysine for glutamate substitution at residue 487 of the mature enzyme (3, 4). The protein encoded by this allele, named ALDH2K for the amino acid substitution, had a markedly increased $K_m$ for NAD$^+$ and reduced $V_{max}$ when expressed as homotetramers in a bacterial expression system (5). This change in $K_m$ for NAD$^+$ renders the enzyme nearly inactive at the concentration of NAD$^+$ that occurs in cells. Moreover, heterozygous individuals, in whom various heterotetramers of ALDH2K and normal ALDH2E subunits are present, are deficient in ALDH2 activity as well (6). The dominance is not complete, as heterozygotes have a milder flush reaction than ALDH2*2 homozygotes (7).

This dominant effect was confirmed recently by this laboratory using retrovirally transduced cell lines expressing one or both alleles. Introduction of the ALDH2*2 cDNA into cells expressing ALDH2*1 resulted in a reduction in enzyme activity. It was assumed that the subunits associated randomly. The degree of reduction in activity in cell lines expressing varying ratios of the ALDH2E and ALDH2K subunits was fit to a model in which only ALDH2E tetramers and (ALDH2E)$_3$-(ALDH2K)$_1$ heterotetramers were enzymatically active, with the normal tetramer having two active sites per subunit and the heterotetramer having one active site (6). Other tetramers containing more ALDH2K subunits were apparently inactive under the assay conditions used. This model accounted for a substantial dominant effect of ALDH2*2, predicting that the heterozygotes would have $\sim$18% of the normal activity in liver of homoygotes.

However, this model did not account for the absence of ALDH2 activity observed with liver extracts from ALDH2*2 heterozygotes analyzed by electrophoresis and staining for enzyme activity. Therefore, additional mechanisms to explain the low ALDH activity were sought. It was noted that in the cells expressing both ALDH2 alleles, the total amount of ALDH2 protein detected by quantitative Western blots was lower than expected (ranging from 1.2 to 2 times that of the cells expressing only ALDH2E from which the doubly transduced cells were derived) (6). In addition, studies performed before the genetic basis for the deficiency was understood suggested that liver extracts from individuals deficient in ALDH2 activity had less immunologically cross-reactive ALDH2 protein than individuals with active ALDH2 (8, 9). Preliminary measures of the enzymes’ half-lives in the transduced HeLa cells suggested no difference between ALDH2E and ALDH2K (6). However, the method used tends to underestimate the half-life of metabolically stable proteins. Therefore, we have extended the preliminary observations by Western blotting liver extracts to determine ALDH2 protein levels and by measuring the turnover of the enzyme in the ALDH2-expressing cell lines by a combination of approaches. We now report a dominant effect of the ALDH2K subunit on the turnover of the enzyme.
Methods

Most chemicals were from Sigma Chemical Co. (St. Louis, MO). Antibodies directed against 3-hydroxyisobutyrate dehydrogenase were kindly provided by Dr. Robert Harris (Indiana University). Radioactively labeled cysteine and methionine ([35S]-Translabel) and [32P]-labeled protein A were from ICN Biochemicals, Inc. (Irvine, CA). Liver samples were provided by Dr. T.-K. Li and Dr. William Bosron (Indiana University) and were genotyped at the ALDH2 locus (10) by the Molecular Biology Core of the Indiana Alcohol Research Center.

Cell lines and culture conditions. The retrovirally transduced cells used for this study were described previously (6). They were maintained in minimal essential medium with 5% fetal calf serum, 100 μg streptomycin/ml, and 63 μg penicillin G/ml. For determination of protein degradation rates, the cells were split and grown overnight. In the morning, penicillin (100 μg/ml) was added. The zero time cells were harvested 30 min later, and cells were subsequently harvested 4, 8, 16, 20, and 24 h later. The cells were rinsed, scraped into 1 ml of phosphate-buffered saline, and centrifuged at 14,000 × g for 2 min. The cell pellets were frozen at −80°C until analysis. Cell proteins were extracted by suspending them in 50 mM sodium phosphate, pH 6.0, 1% Triton X-100, 1 mM PMSF, 1 mM dithiothreitol, and 1 mM EDTA, then briefly sonicating them on ice. The suspension was then centrifuged for 10 min at 14,000 × g and the supernatant was used for quantitative Western blotting as described previously (6). Standard curves had been shown previously to be linear in this range of protein loading.

Analysis of ALDH2 levels in human liver samples. Autopsy liver samples were stored at −80°C until use. 100-μg samples of frozen liver were homogenized in a Polytron homogenizer in the cell extraction buffer. The homogenate was then centrifuged at 100,000 × g for 1 h. The supernatants were analyzed by isoelectric focusing and Western blotting as described (6). As an internal control for degradation of proteins, a separate aliquot of extract was fractionated by SDS-PAGE and Western blotted with antiserum raised against 3-hydroxyisobutyrate dehydrogenase. The antibody bound to the proteins was visualized by incubating the blot with [32P]-labeled protein A, followed by washing and autoradiography. The blots were quantified by β-scanning the filters. The radioactivity in the bands was corrected for background by subtracting the average radioactivity in the same area of the filter above and below the band.

Pulse–chase labeling. Cells were replated 1 d before the experiment and used the following morning at a density of 1.5 × 106 cells per 60-mm dish. The cells were first incubated in cysteine- and methionine-free medium to deplete intracellular amino acid pools. They were then incubated with 40 μCi of [35S]-Translabel (a mixture of [35S]-labeled methionine and cysteine derived from Escherichia coli lysates) in 0.4 ml of cysteine- and methionine-free medium for various times. Biosynthetic rates were determined by harvesting the cells at 1, 2, 4, and 8 h and determining the incorporation of label into immunoprecipitable ALDH2. Degradation rates were determined by labeling the cells for 8 h, then rinsing the cells repeatedly to remove the labeled amino acids, and culturing them for varying times in standard medium. Immunoprecipitations were carried out by incubating 20 μg of cell extract (prepared as above) with 10 μl of preimmune serum for 2 h at 4°C. Nonspecifically bound proteins were removed with 20 μl of protein A Fast Flow-4. Supernatants were then incubated with 2 μl of purified anti-ALDH2 antibody for 8 h. The immune complexes were then recovered by adding 20 μl of protein A FastFlow-4 and incubating an additional 1 h. The complexes were pelleted at 14,000 g and were washed three times with 1 ml of the immunoprecipitation buffer. The entire pellet was then denatured and the supernatant was analyzed by SDS-PAGE (11). The gels were dried and the radioactivity in the bands was quantified by β-scanning.

Statistical analysis. Differences in the amounts of ALDH2 in livers of different genotypes as well as in rates of protein synthesis or degradation were determined by ANOVA (SPSS Inc., Chicago, IL). Degradation and synthetic rates were determined by least-squares fits of the logarithm of counts per minute in the ALDH2 bands plotted against time (12), using Microsoft Excel.

Results

Levels of ALDH2 immunoreactivity in human liver samples. Polyclonal antibodies were generated against purified bovine liver ALDH2. The antibodies were characterized by Western blotting different amounts of extracts of ALDH2E- and ALDH2K-expressing cells, developing the blot with the antibodies and radiolabeled protein A, and quantifying the radioactivity in the bands with a β-scanner (similar to the first three lanes of Fig. 1A). The slopes of the plots of radioactivity per band versus the amount of protein loaded were the same for ALDH2E and ALDH2K (not shown), indicating that the antibodies detected ALDH2E and ALDH2K proteins equally well. 13 human liver samples were genotyped at the ALDH2 locus, and protein extracts were analyzed by quantitative Western blotting of isoelectric focusing gels. Fig. 1A shows a representative Western blot. Equal amounts of protein were loaded for each individual. The isoelectric points of the proteins confirmed the genotypes in each case. The intensity of the bands suggested that the level of ALDH2 immunoreactive material was higher in the ALDH2*1 homozygotes than in heterozygotes. Since only a single ALDH2*2 homozygous
liver was available for analysis, it was impossible to determine differences in enzyme protein levels between ALDH2*2 hetero- and homozygotes.

A possible explanation for the lower amount of ALDH2 seen in the heterozygotes was random differences in postmortem degradation of proteins. A control for this possibility was determination of the level of another mitochondrial matrix space protein. Therefore, Western blots were performed to assess the levels of 3-hydroxyisobutyrate dehydrogenase, a matrix space enzyme involved in valine catabolism (Fig. 1 B). These blots showed relatively constant levels of this protein from liver to liver. The Western blots were quantified using a β-scanner. The radioactivity in the ALDH2*1 homozygote extracts was 540±60 cpm/band (means±SE, n = 9), while that in the ALDH2*2 group was 240±10 cpm/band (n = 4, P < 0.02). The level of 3-hydroxyisobutyrate dehydrogenase was 490±40 and 450±70 cpm/band, respectively (P = 0.64). The ratios of ALDH2 to HIBA were 1.2±0.2 and 0.6±0.1 (means±SE) for the normal versus ALDH2-deficient groups. Thus, livers from ALDH2*2 hetero- or homozygotes had about half the ALDH2 protein level of the livers with active ALDH2.

**Turnover of ALDH2 isozymes in transduced cell lines.** There are no hepatoma or other cell lines that express ALDH2*2; therefore, cell lines expressing one or both ALDH2 alleles previously constructed in the laboratory were used (6). In addition to cells transduced with only ALDH2*1 or ALDH2*2 expressing vectors, one cell line (HeLa EK12) was selected for further study. It represents the situation predicted in heterozygous liver cells, since it expressed equal levels of ALDH2*1 and ALDH2*2 transcripts. These cells were shown previously to form heteromeric enzymes with isoelectric points between those of ALDH2E and ALDH2K (6). ALDH2 protein was found at the highest level in the mitochondrial fraction of these cells prepared by sucrose gradient centrifugation and by immunocytochemistry (data not shown) and the ALDH2E and ALDH2K enzymes had the same pI as the liver enzymes, indicating that the ALDH2 subunits are correctly transported into and processed in the mitochondria. Thus, they are a reasonable model for the study of ALDH2 turnover.

The first approach taken was to follow the level of ALDH2 protein in cells treated with puromycin. In the absence of new protein synthesis, the decline in protein levels reflects the rate of degradation. The cells were cultured for 4–24 h after the addition of puromycin at a concentration reported to completely inhibit protein synthesis (13), and extracts of the cells were Western blotted to quantify the level of ALDH2 protein remaining (Fig. 2). Longer periods of incubation were not possible due to the toxicity of puromycin. Originally, we had reported that the half-life of ALDH2E was 26 h (6). With additional experiments, the estimated half-life was 53±21 h (n = 6 replicate experiments). The large standard deviation reflects the inaccuracy in determining the small changes in protein level over 24 h for a stable protein, since the fitting of data is disproportionately affected by the 24 h time point. On the other hand, the half-life of the ALDH2K-containing proteins was 16±3 h (n = 5) for cells containing heterotetramers, and 19±6 h (n = 6) for cells containing homotetrameric ALDH2K (the latter similar to our initial estimate of 24 h [6]). These two half-lives were significantly less than that for ALDH2E (P < 0.005), but were not significantly different from each other.

To obviate the problems inherent in inhibitor studies, the biosynthetic rate and half-life of the proteins were then determined by pulse–chase experiments. Results of a representative experiment are shown in Fig. 3 A. During the pulse phase, the rate of incorporation of label into immunoprecipitable ALDH2 was not significantly different for the three cell lines. During the chase with unlabeled amino acids, the rate of loss of radioactivity was greater for ALDH2*2-expressing cells (HeLa CK and HeLa EK) than in the cells expressing only ALDH2*1 (Fig. 3 B). The half-lives estimated from this method were 22±8 h for ALDH2E (n = 5), 14±5 h for ALDH2K (n = 3), and 13±4 h for cells expressing both subunits (n = 5, P < 0.05). These results are summarized in Table I.

![Figure 2. Rate of decay of ALDH2 protein after treatment of cells with puromycin.](image)

**Figure 2.** Rate of decay of ALDH2 protein after treatment of cells with puromycin. HeLa CE (E cells), HeLa CK (K cells), or HeLa EK (EK cells) cell lines were treated with puromycin, then harvested at the different time points shown. Cell extracts were analyzed for ALDH2 content by quantitative SDS-PAGE Western blotting and autoradiography.

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Protein half-life (h)</th>
<th>Synthetic rate (cpm/h) pulse label</th>
</tr>
</thead>
<tbody>
<tr>
<td>HeLa CE</td>
<td>53±21 (6)</td>
<td>22±8 (5)</td>
</tr>
<tr>
<td>HeLa EK</td>
<td>16±3 (5)*</td>
<td>13±4 (5)*</td>
</tr>
<tr>
<td>HeLa CK</td>
<td>19±6 (6)*</td>
<td>14±5 (3)*</td>
</tr>
</tbody>
</table>

The cells lines were clonal lines expressing ALDH2E (HeLa CE), ALDH2K (HeLa CK), or both subunits (HeLa EK) (6). Decay in protein mass after puromycin was determined by Western blotting. Decay in protein mass in the pulse–chase experiments was determined from loss of immunoprecipitable labeled ALDH2, and synthetic rate was determined as the increase in immunoprecipitable labeled ALDH2. The number of separate experiments is shown in parentheses, and the data given are means±standard deviation. *Statistically significant differences from the HeLa CE cells by ANOVA (P < 0.05).
Effect of increased turnover of ALDH2K on expression of ALDH2 activity in heterozygotes. The effect of reduction in half-life on the residual activity of ALDH2 can be estimated from the amount of ALDH2K-containing tetramers predicted to be formed in heterozygous cells (Table II). Each ALDH2 allele directs synthesis of $n$ subunits; thus, there are $2n/4$ or $n/2$ tetramers. If subunits assemble randomly, $6\%$ ($1/16$) of the tetramers will be ALDH2E homotetramers with normal half-life. The remaining tetramers contain at least one ALDH2K subunit. The model that best accounted for the residual activity in cells expressing both alleles permitted two active sites for E$_4$ and one active site for E$_3$K tetramers, predicting 37% residual activity (6). Table II shows the original model and corrections for the shorter half-life of heterotetramers (assumed here to result in a 50% reduction in protein levels). These models are compared with the activity expected for the parental cells expressing $n/2$ active sites of ALDH2E (Table II). The reduced level of E$_3$K tetramers reduces the activity from 37 to 25%, in reasonable agreement with observed residual activity (31% of that in the ALDH2E-expressing cells from which the doubly transduced cells were derived [6]). If the reduction in E$_3$K subunits is 34% (based on the direct measurements in the HeLa cells), the agreement is even better (29% activity predicted). The residual activity in heterozygous livers would be 12% of normal livers, since the acquisition of the ALDH2*2 allele is accompanied by the loss of one of the ALDH2*1 alleles. This is in the range of residual low $K_m$ ALDH activity reported in deficient human livers (7, 14).

### Discussion

Human ALDH2 deficiency is an interesting genetic trait. From a clinical standpoint, it is the single strongest genetic factor that influences drinking behavior and risk of alcoholism (15–

**Table II. Models for the Dominance of the ALDH2*2 Allele**

<table>
<thead>
<tr>
<th>Enzyme tetramer</th>
<th>Fraction of total</th>
<th>Active sites per tetramer</th>
<th>Uncorrected active sites</th>
<th>Correction for half-life</th>
<th>Corrected active sites</th>
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</thead>
<tbody>
<tr>
<td>E$_4$</td>
<td>1/16</td>
<td>2</td>
<td>$n/16$</td>
<td>100%</td>
<td>$n/16$</td>
</tr>
<tr>
<td>E$_3$K</td>
<td>4/16</td>
<td>1</td>
<td>$2n/16$</td>
<td>50%</td>
<td>$n/16$</td>
</tr>
<tr>
<td>E$_2$K</td>
<td>6/16</td>
<td>0</td>
<td>0</td>
<td>50%</td>
<td>0</td>
</tr>
<tr>
<td>EK$_3$</td>
<td>4/16</td>
<td>0</td>
<td>0</td>
<td>50%</td>
<td>0</td>
</tr>
<tr>
<td>K$_4$</td>
<td>1/16</td>
<td>0</td>
<td>0</td>
<td>50%</td>
<td>0</td>
</tr>
</tbody>
</table>

The calculations are based on the model presented previously (6) for ALDH2 heterotetramers. This model permits two active sites for E$_4$ and one active site for E$_3$K tetramers and assumes synthesis of $n$ subunits per allele; thus, there will be $2n/4$ ($= n/2$) tetramers synthesized. This value is multiplied by the fraction of each tetramer and the number of active sites per tetramer to obtain the uncorrected number of active sites. The corrected number of active sites takes into account the increased turnover of the E$_3$K tetramers. For comparison, the parental cells expressing only one ALDH2*1 allele will synthesize $n$ subunits, $n/4$ tetramers, and therefore $n/2$ active sites. Without the correction for turnover of the heterotetramer, the activity would be 37% of that of the parental cells ($3n/16$ divided by $n/2$), and with the correction, 25% of the activity of the parental cells ($n/8$ divided by $n/2$).

2030  Xiao et al.
ALDH2 protein in ALDH2-deficient livers (8, 9) by showing that ALDH2K-containing enzymes are degraded more rapidly than ALDH2E enzymes. The difference in half-lives between ALDH2E and ALDH2K enzymes determined here would result in a 34% reduction in total ALDH2 protein [taking into consideration the normal half-life of the small fraction of (ALDH2E)4 tetramers]. The difference between this value and the 50% reduction in ALDH2 in the liver extracts may simply reflect the well-known tissue-specific differences in enzyme degradation rates for the same protein (18). The mechanism by which ALDH2K exerts this effect is unknown. No difference in half-life of the two enzyme forms was apparent in E. coli expressing these proteins (5), suggesting that the difference relates to the subcellular compartmentation of ALDH2. Mitochondrial proteins are assisted in folding and in degradation by a family of heat shock proteins (19–22) and probably by a protease (PIM) characterized in yeast mitochondria (23). The ALDH2K subunit may slow the assembly of the tetramer and perhaps render it prone to more rapid catabolism. Such a mechanism has been demonstrated for medium chain acyl-CoA dehydrogenase mutants (24). Alternatively, the reduced affinity of the enzyme for NAD+ may contribute to its higher turnover. The inhibition constant for NAD+ (Km, equivalent to the dissociation constant) for ALDH2E enzyme is 13 µM, whereas for the ALDH2K enzyme it is ~500 µM, due to a much reduced rate of association of enzyme with coenzyme (5). Thus, at the estimated intramitochondrial-free NAD+ concentration of 6 mM (25), virtually all the wild-type enzyme would contain bound NAD+, whereas ~8% of the ALDH2K enzyme would be coenzyme free, and potentially more liable to degradation. ALDH2 heterotetramers synthesized in E. coli had only a single, rather low Km for NAD+, but subunits without bound NAD+ probably still are present in the heterotetramers (Wang, X., S. Sheikh, D. Saigal, L. Robinson, and H. Weiner, manuscript submitted for publication).

Increased turnover of mutant enzymes has been observed in numerous genetic diseases. In some cases the mutant enzyme is stabilized and enzyme activity is increased by supplementation of the diet with a vitamin precursor of the enzyme cofactor (for example, riboflavin for flavoproteins [26], pyridoxine for pyridoxal enzymes, and thiamine for thiamine pyrophosphate–dependent enzymes). These genetic deficiencies have invariably been recessively inherited. A possible exception is an unstable short chain acyl-CoA dehydrogenase in a compound heterozygote (27, 28), but it is not known if one allele was acting dominantly or if both alleles contributed to the instability. Certain porphyrias are dominantly inherited and three involve intramitochondrial enzymes, but these are examples in which a 50% reduction in enzyme activity is sufficient to cause the clinical phenotype, and the mutations are not dominant negatives. The ALDH2*2 allele appears to be the first example of a dominantly acting mutation that alters turnover of a mitochondrial protein. It will be interesting to test whether manipulating mitochondrial NAD+ levels modifies the half-life of the ALDH2K enzymes.

Acknowledgments

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References