In Vivo Effects of Protease Inhibitors on Chickens with Hereditary Muscular Dystrophy

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ABSTRACT Beginning on day 4 ex ovo, and every 3 d thereafter, genetically dystrophic Line 413 chickens were given intraperitoneal injections (4 mg/kg body wt) of a protease inhibitor, leupeptin, pepstatin, or antipain. Experimental chickens received protease inhibitors dissolved in a water:ethanol:dimethylsulfoxide solution (50:40:10, vol:vol:vol). Control untreated animals received diluent injections.

Untreated dystrophic chickens typically reach around day 30 ex ovo a maximum ability to right from the supine position in a standardized functional test for muscle weakness. After day 30 ex ovo, the dystrophic chickens are found to decline progressively in their ability to right, compared with normal, nondystrophic controls, which have an unimpaired ability to right. Concomitantly, dystrophic chickens exhibit characteristically high levels of plasma creatine phosphokinase enzyme activity. In addition, an increased frequency of degenerating, regenerating, and vacuolated myofibers, and inflammatory cells appear in the affected pectoralis major muscles from the dystrophic chicken.

 Throughout the duration of the trial, there was no major enhancement in the functional righting ability of dystrophic chickens receiving any one of the protease inhibitors tested. However, there was a significant reduction in the abnormally high levels of plasma creatine phosphokinase in the treated chickens. Also, there was an apparent reduction in the mean number of vacuolated fibers in the pectoralis muscle from the protease inhibitor-treated birds. No significant reductions were observed in the relative frequency of degenerating and regenerating myofibers or inflammatory cells. In addition to the plasma creatine phosphokinase decrease, however, therapeutic benefit was seen in 31.0, 30.5, and 14.8% increases in the wet weight (and total noncollagen protein) of pectoralis muscle from dystrophic chickens receiving leupeptin, pepstatin or antipain, respectively.

INTRODUCTION

Substantial elevation in proteolytic enzyme activity is associated with the progressive loss in muscle tissue exhibited in both genetically dystrophic humans (1-3) and animals (4-7). Since the pioneering biochemical studies of Umezawa and coworkers (8, 9), in which microbial inhibitors to proteolytic activity were isolated, evidence has indicated that some of these inhibitors (e.g., leupeptin, pepstatin, and antipain) may have a therapeutic role in retarding the abnormal muscle protein degradation found in hereditary myopathies. In this regard, the addition of protease inhibitors in vitro to embryonic, intact, or homogenate muscle preparations from dystrophic animals promotes increased protein synthesis (10), decreased protein breakdown (10-13), and decreased atrophy in general (14).

There has been a lack of trials in vivo involving protease inhibitor treatment of animals with genetic myopathies. Results to date are inconclusive; for example, when dystrophic mice were given either subcutaneous or intraperitoneal dosages of protease inhibitors, benefit was seen in one study (15), but not in another (16). In another report (17), improvements in muscle mass and histology and significant reductions in abnormally high levels of serum creatine phosphokinase activity (CPK)1 were found in genetically dystrophic chickens injected with leupeptin and pepstatin intramuscularly.

The purpose of the present study was to investigate further the effects in vivo of individual protease inhibitors on the genetically dystrophic Line 413 chicken within an established, standardized, drug screening program (18-21). Reported here is new evidence for limited degrees of drug-related improvements in a

1Abbreviations used in this paper: CPK, creatine phosphokinase activity; FN, flip number.
functional test for righting ability and plasma CPK levels, and in muscle mass and histology.

METHODS

Line 413 dystrophic and genetically related Line 412 normal, nondystrophic chickens were purchased as eggs from the Department of Avian Sciences, University of California, Davis. The incubation conditions and subsequent care and maintenance of the mostly female chicks were as previously described (18–21).

Starting on day 4 ex ovo, each experimental chicken received an intraperitoneal injection of 0.5 ml protease inhibitor solution (4 mg/kg body wt) every 3 d between 0800–1000 h. Each protease inhibitor was dissolved in a water:ethanol:dimethyl sulfoxide solution (50:40:10, vol:vol:vol). A number of dystrophic chickens in each trial received diluent only and served as untreated dystrophic controls. All animals were weighed twice a week.

The functional assessment of muscle strength was measured with the extensively described flip test procedure (18–21). In brief, each drug-treated and untreated control chicken was given a standardized test for righting from the supine position every 5 d between 1300 and 1500 h. The number of successful attempts to regain an upright position scored out of five consecutive opportunities was defined as the flip number (FN). To eliminate bias in testing, all chickens were pooled before the testing procedure. Subsequently, each bird was randomly selected for the righting test, and the FN later matched with its wing band number.

Blood for the determination of plasma CPK was obtained without anesthesia from the jugular vein into heparinized syringes between 1000 and 1200 h. Plasma was separated by centrifugation at 755 g for 15 min at 4°C. On the same day on coded samples, plasma CPK activities were assayed by an ultraviolet spectrophotometric procedure, using the activated CPK reagent set of Boehringer Mannheim Biochemicals, Indianapolis, Ind. (cat. No. 124176). Enzyme activity was expressed in International milliunits CPK per milliliter plasma at 25°C, pH 7.0. Plasma dilutions were made when enzyme activities were near or above the linear range of the assay conditions.

To test the possible inhibitory effect of the protease inhibitors directly on CPK, aliquots of plasma with known CPK activity were assayed individually with the addition of 0.081 mM leupeptin, 0.054 mM pepstatin, or 0.062 mM antipain. The water:ethanol:dimethyl sulfoxide diluent (50:40:10; vol:vol:vol) of the protease inhibitors was also tested at the same final concentration of 1.8% for its effect on the assay of CPK. As a result, neither the protease inhibitors nor the diluent was found to have an effect on the quantitation in vitro of plasma CPK activities.

In the histological survey of muscle involved in the righting reflex, samples of pectoralis major muscle from each chicken were removed at day 67 ex ovo after completion of the drug trials. As previously described (20), each muscle sample was excised, weighed, and placed in an isometric clamp before immersion in 4% paraformaldehyde, fixed for 18–24 h, and embedded in paraffin. 4–6-μm sections were stained with hematoxylin, eosin, and Masson’s trichrome. Four pathologic criteria were routinely examined and quantitated in muscle specimens from each bird: vacuolation, fiber necrosis, inflammation, and fiber regeneration. Employing a calibrated grid, 16 individual microscopic fields per muscle were examined and each of the above pathologic features was enumerated. The frequencies of necrotic and regenerative fibers were grouped together. Values were expressed as the mean number of observations per square millimeter.

Total noncollagen protein of the pectoralis major muscle from the experimental chickens was determined in homogenates by the procedure of Lowry et al. (22), using bovine serum albumin as standard.

Leupeptin, pepstatin, and antipain were donated by Dr. Walter Troll, New York University School of Medicine, New York, Dr. Arthur M. Dannenberg, Department of Environmental Medicine, Johns Hopkins University, Baltimore, Md., and Dr. Hamao Umezawa, Institute of Microbial Chemistry, University of Tokyo, Japan. All chemicals were reagent grade. Statistical tests of significance of paired means were derived from the two-sided Students’ t test.

RESULTS

Flip test for righting ability. Dystrophic chickens treated with leupeptin exhibited a temporary yet significant (P < 0.001) prolongation in a functional test for muscle strength (Fig. 1). As seen here, and has been reported extensively elsewhere (19–21), the untreated dystrophic control Line 413 chickens reached an optimal FN around day 30 ex ovo and subsequently declined. However, leupeptin-treated chickens maintained a perfect FN of five for 2 additional wk before decreasing in righting ability.

There were no significant differences evident in the righting abilities of dystrophic Line 413 chickens treated with either pepstatin (Fig. 2) or antipain (Fig. 3).

![Figure 1: Effect of leupeptin on mean FN of Line 413 dystrophic chickens. The mean FN of the leupeptin-treated group (○) at days 39 and 44 ex ovo are significantly different from the corresponding untreated control group (●) at P < 0.001. Details of dosage schedule and procedure for flip testing are as described in Methods. The number of animals per point in four untreated dystrophic control and six leupeptin-treated dystrophic.](image-url)
FIGURE 2  Effect of pepstatin on mean FN of Line 413 dystrophic chickens. Details of dosage schedule and procedure for flip testing are as described in Methods. The number of animals per point is 5 untreated dystrophic control (●) and 10 pepstatin-treated dystrophic (○).

3) compared with the corresponding untreated dystrophic control chickens. Normal control chickens reached a perfect mean FN of five at day 15 ex ovo and subsequently maintained this FN throughout the duration of the trials (data not shown).

Plasma creatine phosphokinase activity. As seen in Fig. 4, there were significantly (P < 0.001) higher levels of CPK activity in the plasma of dystrophic Line 413 chickens compared with the normally low values exhibited in the nondystrophic Line 412 chickens in each age group. Though CPK values were much higher than normal, significant reductions (P < 0.001) in plasma CPK were elicited in dystrophic chickens treated with leupeptin (Fig. 4, trial A), pepstatin, or antipain (Fig. 4, trial B). Each of these reductions was found throughout the duration of the drug trial.

Quantitation of muscle histology. In pectoralis major muscle from the untreated normal Line 412 chickens, no fiber necrosis, regeneration, vacuolar change, or inflammation was present (Table I). However, pathologic features observed in muscle from untreated dystrophic chickens were quite uniform. Vacuolar change, fiber necrosis, regenerative activity, and inflammatory cells were widespread in all specimens. Vacuolar change in the dystrophic myofiber occurred as multiple, round, clear, unstained sarcoplasm. Necrotic and degenerative fibers were granular and fragmented, and were often accompanied by phagocytic activity. Regenerating fibers, easily recognized in hematoxylin and eosin stains by their blue sarcoplasm (23), typically had multiple, large vesicular nuclei with prominent nucleoli. Inflammatory infiltrates found in the dystrophic muscle were primarily perivascular. They were composed of a monotonous, mononuclear
cellular population of lymphoid cells having small, dark nuclei and little cytoplasm.

No significant differences were found in the relative frequencies of regenerating and degenerating fibers and inflammatory cells between muscle samples from untreated dystrophic controls and chickens receiving protease inhibitor treatment (Table I). However, there were significant reductions ($P < 0.01$) in the mean number of vacuolated fibers found in a square millimeter of muscle from pepstatin- or antipain-treated chickens, compared with the untreated dystrophic controls (Table I).

**Body and tissue weights.** Little if any change in body weight was found in the main female dystrophic chickens treated with the various protease inhibitors compared with the corresponding untreated female dystrophic controls. Significant increases of 31.0, 30.5, and 14.8% were found in the wet weight of the afflicted pectoralis major muscle (expressed as percent body weight) from leupeptin-, pepstatin-, and antipain-treated dystrophic chickens, respectively. In this regard, there was no significant difference found in noncollagen protein of dystrophic muscle (mean 101.5 ± 22.0 mg noncollagen protein/g wet wt muscle) compared with muscle from chickens treated with any of the protease inhibitors (95.5±17.5 mg/g). However, it should be noted that the muscle noncollagen protein from either the drug-treated or untreated dystrophic chickens was found to be significantly less ($P < 0.001$) than that measured in comparable normal chicken muscle (227.3±17.2 mg/g).

Compared with the untreated dystrophic controls, small increases (4.0, 12.1, and 5.4%) in the liver wet weight were found in dystrophic chickens treated with leupeptin, pepstatin, and antipain, respectively. No change in the heart tissue was observed with any of the drug therapies.

**DISCUSSION**

In contrast to the significant enhancement in righting ability exhibited in Line 413 dystrophic chickens treated with either antiserotoninergic drugs (20, 21) or diphenylhydantoin (24–26), no remarkable improvement was demonstrated with protease inhibitor treatment. The temporal benefit in righting ability derived from leupeptin may be due in part to intraperitoneal leupeptin being able to penetrate muscle tissue better than either pepstatin or antipain (11, 27). However, it should be noted here that the FN remains a standardized measurement of functional ability or disability in the chicken and is not a direct quantitation of inherent muscle strength or weakness (18–21, 24, 25). In future drug trials, a technique from Cabe et al. (28, 29) will be adapted to the chicken to determine the relative strength of the primary wing muscles. They use a sensitive strain gauge to assess directly the maximal strength of rat and mouse hindlimb and forelimb muscles in routine toxicological studies.

The abnormal elevation in blood CPK activity in dystrophic chickens (18–21, 24, 25, 30) and humans (31) is generally accepted as an indirect indicator of the onset and rate of progression of the muscle disease. In this connection, the significant reductions in plasma CPK levels found with each of the protease inhibitors (given intraperitoneally) can be perceived as therapeutically beneficial. This finding compares with previous reports of CPK reductions in dystrophic chickens given either subcutaneous (32) or intramuscular (17) injections of protease inhibitors. However, caution should be used in interpreting the relative value of reduced blood CPK as a criterion of drug-related benefit. For example, Munsat and Bradley (33) found that increases in CPK levels were not accompanied by significant clinical improvement in individuals with facioscapulohumeral dystrophy treated with prednisone.

**TABLE I**

Quantitation of Pectoralis Major Muscle Histology

<table>
<thead>
<tr>
<th>Experimental condition</th>
<th>n</th>
<th>Degenerating and regenerating fibers</th>
<th>Vacuolated fibers</th>
<th>Inflammatory cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal—untreated</td>
<td>5</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Dystrophic—untreated</td>
<td>7</td>
<td>4.7±0.7</td>
<td>15.7±2.0</td>
<td>7.3±2.1</td>
</tr>
<tr>
<td>Dystrophic—leupeptin</td>
<td>6</td>
<td>4.8±0.5*</td>
<td>12.9±1.8*</td>
<td>4.5±0.8*</td>
</tr>
<tr>
<td>Dystrophic—pepstatin</td>
<td>6</td>
<td>3.1±0.7*</td>
<td>5.8±1.3†</td>
<td>4.6±1.3*</td>
</tr>
<tr>
<td>Dystrophic—antipain</td>
<td>5</td>
<td>3.7±0.7*</td>
<td>4.6±1.7†</td>
<td>3.8±1.9*</td>
</tr>
</tbody>
</table>

For drug schedule and histologic procedures see Methods. Day 67 ex ovo birds were sacrificed at the completion of the drug trial. Values are expressed as the mean number of observations±SEM per square millimeter, and are based on the enumeration of sixteen separate 1-mm² grids per muscle sample.

* Difference from the corresponding untreated dystrophic group is not significant.
† Difference from the corresponding untreated dystrophic group is significant at $P < 0.01$. 

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The effect of each protease inhibitor on the dystrophic pectoral muscle is inconclusive. In the present study, no significant differences in the relative frequencies of degenerating and regenerating muscle fibers were noted with treatment. In addition, the relative reduction in the mean number of vacuolated fibers is masked by the variability of these observations among individual muscle specimens, and between treated and untreated animals. To a great degree, these findings do not concur with the improved histological appearance previously reported for pectoralis muscle from dystrophic chickens treated with pepstatin (15). However, the drug-related increases in muscle mass and in noncollagen protein found here are very encouraging and are consistent with the increased myofiber areas observed microscopically in muscles from protease inhibitor-treated dystrophic chickens (17) and mice (15).

The major benefits from this trial of intraperitoneally administered leupeptin, pepstatin, or antipain are significant reductions in plasma CPK and net increases in pectoralis major muscle mass and noncollagen protein. It remains unresolved whether there is a drug-induced enhancement in actual muscle strength capability, which has been assessed indirectly by either the functional test for righting ability, quantitative muscle histology, or myofibrillar protein content (viz., non-collagen protein).

ACKNOWLEDGMENTS
The authors wish to thank Mr. James A. Stamos for his preparation of the illustrations, and Mr. Barry Wolitzky for his technical assistance.

Michael S. Hudecki holds a Research Career Development Award from the National Institute of Neurologic Disease and Stroke (NINDS). This work was supported by grants from the Muscular Dystrophy Association and NINDS.

REFERENCES