THE EFFECT OF SPONTANEOUS ELECTROMECHANICAL ACTIVITY ON THE METABOLISM OF ACETYLCHOLINESTERASE IN CULTURED EMBRYONIC RAT MYOTUBES

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Abstract

We have investigated the effect of electromechanical activity on the molecular forms of acetylcholinesterase (AChE) in cultured embryonic rat myotubes. Both globular and asymmetric forms of AChE are present on the 5th day of culture when myotubes are just beginning to fibrillate. Between days 5 and 8, the 4 S (G1), 10 S (G2), and 16 S (A12) forms increase dramatically, and appreciable 12.5 S (A8) AChE appears. When fibrillation is prevented by adding tetrodotoxin on day 4, the increases in the A12 and A8 forms are prevented, and the increases in the G1 and G2 forms are significantly impaired. At 8 days, fibrillating myotubes have 19 times more A12 AChE and over 4 times more G1 and G2 enzyme than do nonfibrillating myotubes. The effect of tetrodotoxin is reversible. When tetrodotoxin is removed at 7 days, fibrillation resumes promptly, and globular and asymmetric forms recover. Light microscopic examination of fibrillating and nonfibrillating myotubes showed that tetrodotoxin does not affect the gross morphological development of the myotubes. Titration of AChE-active sites with O-ethyl-5,5-diisopropyl methyl-phosphonothionate demonstrated that the increase in AChE activity associated with fibrillation is due to an increase in the number of AChE molecules present and not to an increase in the rate at which individual AChE molecules turn over acetylcholine. To evaluate AChE metabolism in fibrillating and nonfibrillating myotubes, we examined the enzyme after inactivating it with paraoxon. Paraoxon readily penetrates cells and diethylphosphorylates a serine in the active site of AChE, thereby inactivating it. The diethylphosphorylated enzyme is stable, but it can be reactivated rapidly and quantitatively with pyridine-2-aldoxime methiodide (2-PAM). After inactivating AChE with paraoxon, we simultaneously evaluated synthesis (by following the newly synthesized active AChE) and turnover (by following the 2-PAM-reactivatable AChE). Our results show that globular and asymmetric forms of AChE are both synthesized more rapidly in fibrillating than in nonfibrillating myotubes.

A central concern of cell biology is the regulation of cell surface macromolecules. One cell surface protein that is of considerable interest is the enzyme acetylcholinesterase (AChE). There are at least six molecular forms of this enzyme. Three of these forms are globular proteins and three are asymmetric (Bon et al., 1979). The globular forms of the enzyme are monomers, dimers, or tetramers of identical catalytic subunits designated G1, G2, and G4. The asymmetric forms are designed A4, A8, and A12 and consist of one, two, or three tetramers attached to an elongated collagen-like tail. Muscle cells externalize three types of AChE: globular forms that are secreted, globular forms that are integral sarcolemmal proteins, and asymmetric forms that appear to associate electrostatically with the basal lamina that surrounds each muscle fiber (Rotundo and Fambrough, 1980; Younkin et al., 1982). In all cells that have been studied, a large percentage of total AChE is intracellular (Lazar and Vigny, 1980; Rotundo and Fambrough, 1980; Inestrosa et al., 1981; Taylor et al., 1981; Brockman et al., 1982). Recent studies by Rotundo and Fambrough (1980) on cultured embryonic chick myotubes indicate that this intracellular AChE is newly synthesized enzyme that is destined for externalization.

Most of the AChE in innervated adult mammalian
skeletal muscle is intracellular G1 and sarcolemmal G4 enzyme that is located outside neuromuscular junctions (nonjunctional AChE), but a significant fraction (26% in rat diaphragm) is highly concentrated at neuromuscular junctions with its active sites exposed (Younkin et al., 1982). This junctional AChE is the functionally important enzyme in skeletal muscle that rapidly hydrolyzes the acetylcholine released from motoneuron terminals thereby terminating neuromuscular transmission. Asymmetric forms are of particular interest because they comprise virtually all (84% in adult rat diaphragm) of the functionally important enzyme (Younkin et al., 1982).

The interaction between the α-motoneuron and the skeletal muscle fiber that it innervates is one of the most intensively studied examples of mammalian cell-cell interaction. α-Motoneurons profoundly influence both junctional (A12 and A6) and nonjunctional (intracellular G1 and sarcolemmal G4) AChE, and there is evidence suggesting that innervation also influences the enzyme that is secreted by muscle cells (Inestrosa et al., 1977; Carter and Brimijoin, 1981). Thus this system is a good one for inquiring generally into the mechanisms that are involved in the regulation of cell surface proteins and muscle. In particular, it involves the mechanisms by which neurons regulate the functionally important surface proteins of the cells that they innervate.

The influence of the α-motoneuron on skeletal muscle AChE must begin with one or several signals that pass between the α-motoneuron and the skeletal muscle fiber. These signals must (through what may be a very complicated series of cellular events) alter the metabolism of the enzyme in a way that generates the levels and cellular localization of the various forms of the enzyme found in adult innervated muscle. There is good evidence that the electromechanical activity (action potentials and/or contraction) set up in muscle by α-motoneurons is one important signal that mediates the influence of α-motoneurons on both junctional (Lomo and Slater, 1980; Rubin et al., 1980) and nonjunctional (Drachman, 1972) AChE in skeletal muscle fibers. The electromechanical activity set up in muscle by nerve also profoundly influences the metabolism of functionally important surface proteins in the cells that they innervate.

The aim of this study was to examine the influence of electromechanical activity on AChE metabolism, focusing particularly on the influence of activity on the rate at which the various forms of the enzyme are synthesized. We have carried out this study in cultured embryonic rat myotubes. These cells are not innervated and have no endplates, but they provide a very useful model system in which to study the effect of electromechanical activity on AChE metabolism. Myotubes derived from 18- to 20-day rat embryos contain both globular and asymmetric forms of AChE (Koenig and Vigny, 1978). The myotubes undergo spontaneous electromechanical activity (fibrillation) that can be inhibited conveniently with tetrodotoxin (TTX), a drug that specifically inactivates the voltage-dependent sodium channels in the sarcolemma that are required for action potential generation.
virtually all of the AChE in these cultures is associated with the myotubes.

**Sequential extraction of globular, asymmetric, and non-extractable AChE.** Extraction was carried out using a modification of the method described by Youngkin et al. (1982). Globular forms of AChE were extracted initially (S1) from washed cells using low ionic strength buffer (LIB)—a 10 mM phosphate buffer, pH 7, containing 1% Triton X-100, 5 mM N-ethylmaleimide, 2 mM benzamidine, and 10 mM EGTA. The pellet remaining after the initial extraction was extracted once more with LIB to remove residual globular forms (S2). Asymmetric forms (S3) then were obtained by extracting once with high ionic strength buffer (HIB), a buffer identical to LIB except that it contained additionally 1.0 mM NaCl. The pellet remaining after these three extractions was homogenized in HIB, and the homogenate (H4) was assayed to evaluate the AChE that was not extracted. Previous work has shown that this procedure extracts 92 ± 4% of the activity present in the initial homogenate (Brockman et al., 1982). N-Ethylmaleimide, benzamidine, and EGTA were included in both extraction buffers to inhibit proteolytic degradation of AChE (Youngkin et al., 1982). All homogenizations were carried out in 1.25 ml of buffer using a Polytron (Brinkman Instruments) equipped with a Kinematica generator run at a setting of 8 for 10 sec. To improve recovery, residual homogenate was washed from the Polytron by operating it for 5 sec at the same speed using a fresh 1.25-ml volume of buffer. The original homogenate and the wash were pooled prior to centrifugation to give 2.5 ml of final homogenate. All three centrifugations were carried out for 30 min at 39,000 g in a Sorvall centrifuge equipped with an SS/34 rotor.

In several experiments, we evaluated the percentage of enzyme activity due to true AChE by assaying fractions in the presence and absence of 0.1 mM tetraisopropyl pyrophosphoramide (iso-OMPA), a specific inhibitor of butyrylcholinesterase. iso-OMPA was added to the samples immediately before assay and the assays were run for a period of 2 hr. At 5 days, 70% of the globular activity and 40% of the asymmetric activity in both fibrillating and nonfibrillating myotubes were due to true AChE (resistant to iso-OMPA). At 8 days, 56% of the asymmetric activity in nonfibrillating myotubes and 83% of the asymmetric activity in fibrillating myotubes was due to true AChE. At this time, essentially all of the globular enzyme in both fibrillating and nonfibrillating myotubes was due to true AChE. Thus the large increases in globular and asymmetric enzyme activity that occurred in fibrillating myotubes (Figs. 1 and 3) were overwhelmingly due to increases in true AChE.

**Separation of individual forms on sucrose gradients.** Individual forms of AChE were separated on sucrose gradients as described previously by Youngkin et al. (1982). AChE recovery was evaluated by comparing the AChE activity applied with the summed activities of the gradient fractions and was 69 ± 2% (n = 13).

**AChE assay.** AChE was assayed as described previously (Brockman et al., 1982) using a modification of the radiometric assay developed by Johnson and Russell (1975).

**Quantitation of AChE activity.** The sequential extraction data on myocyte AChE were used to calculate the amounts of globular (S1 ± S2), asymmetric (S3), and nonextractable AChE. The activities of individual globular and asymmetric forms were obtained from the gradient profiles by expressing the activity in each gradient fraction as a percentage of the total activity on the gradient and summing the percentages, from minimum to maximum, in each of the peaks corresponding to particular forms.

**Morphological studies.** On days 5 to 8 of culture, myotubes were examined under a phase contrast microscope fitted with an eyepiece with a 10 x 10 grid. Eight fields were examined on each dish. In each field, 30 of the 100 squares were scored on a scale of 0 to 3. A 0 was scored if there was no myocyte observed in the square, a 1 was scored if one-third or less of the square was filled, a 2 was scored if one-third to two-thirds of the square was filled, and 3 was scored if two-thirds or more of the square was filled. These scores were converted to a percentage of the total surface area covered by myotubes. On each day, three to four dishes of both fibrillating and nonfibrillating myotubes were examined.

**Titration of active sites with O-ethyl-S-3-diisopropylaminoethyl methyl-phosphonothionate (MTP).** Homogenates of fibrillating and nonfibrillating myotubes were prepared by homogenizing 5 and 13 dishes (fibrillating and nonfibrillating) of 8-day myotubes in 1 ml of LIB. The homogenate of the fibrillating myotubes was then diluted so that it had the same level of AChE activity as the homogenate from the nonfibrillating myotubes. An identical amount of MTP (estimated to inactivate half of the active sites) was added to the two homogenates and they were assayed periodically to determine the amount of AChE activity remaining. At the concentration of MTP used (7.5 x 10^-10 M to 9.8 x 10^-10 M), a stoichiometric titration is achieved after approximately 120 min.

**AChE metabolism.** The medium from 7-day-old myotube cultures (conditioned medium) was removed and saved. Fresh medium containing 10 μM paraoxon (37°C) was added and the cultures were incubated for 30 min to inactivate (diethylphosphorylate) almost all (96%) of the AChE activity in the cultures. The cultures then were rinsed three times with PBS (37°C) and the original medium was replaced. At various times cells were washed with PBS and harvested. In some experiments, four fractions were obtained by sequential extraction as described above. In others, only two fractions were examined, globular forms (S1) and the homogenate of the pellet left after extraction of globular forms (H2). Extracts were assayed both before and after reactivation of diethylphosphorylated enzyme with pyridine-2-aldoxime methiodide (2-PAM) to determine the amount of enzyme present in the cultures labeled by paraoxon. Reactivation was accomplished by adding 0.2 mM 2-PAM to the sample and allowing it to sit 45 min at room temperature before diluting and assaying. Previous work (Brockman et al., 1982) has shown that this procedure completely reactivates all diethylphosphorylated enzyme.

**Spontaneous reactivation of diethylphosphorylated enzyme.** To evaluate the stability of diethylphosphorylated enzyme, 8-day-old myotubes first were treated with 10.0
µM paraoxon for 30 min. At the same time, cultures were treated with 1.0 mM 2,4-dinitrophenol and 100 µg/ml of cycloheximide to inhibit cellular metabolism. The cultures then were washed three times with PBS and medium containing 1.0 mM 2,4-dinitrophenol and 100 µg/ml of cycloheximide was added. Cultures were harvested immediately after treatment with paraoxon and 2 hr after treatment. Homogenates were prepared in LIB and AChE activity was determined.

Materials. Rats were from Zivic-Miller Co. (Allison Park, PA). Dulbecco's modified Eagle's medium was from Flow Laboratories (McCLean, VA), fetal calf serum was from K.C. Biological, Hanks' balanced salt solution and antibiotics were from M.A. Bioproducts, collagenase (CLS II) was from Worthington Biochemical Corp. (Freehold, NJ) [3H]acetylcholine was from New England Nuclear (Boston, MA), iso-amyl alcohol was from Fisher, 4a20 cocktail was from Research Products International, (Elk Grove Village, IL), N-ethylmaleimide was from Eastman Kodak Co. (Rochester, NY) and pyridine-2-zamidine, iso-OMPA, sucrose (grade 1), and Triton X-100 were from Sigma Chemical Co. (St. Louis, MO).

Results

Development of the molecular forms of AChE. The results described below were obtained using myotube cultures derived from 20-day rat embryos. The myotubes in these cultures develop as described in earlier studies (Heinemann et al., 1977; Brockman et al., 1982; Younkin et al., 1982) to obtain fractions containing globular, asymmetric, and nonextractable AChE. Figure 1 shows that the enzyme activity in each fraction increased much more rapidly in fibrillating than in nonfibrillating cultures. In both fibrillating and nonfibrillating cultures, globular forms of AChE increased steadily between days 5 and 9 (Fig. 1A), but the increase was greater in fibrillating myotubes (12-fold) than in TTX-treated myotubes (3-fold). Low levels of asymmetric and nonextractable AChE were detectable in nonfibrillating cultures, but the dramatic increases in asymmetric and nonextractable AChE that occurred in fibrillating cultures (14-fold and 3-fold, respectively, between days 5 and 9) did not occur when spontaneous electromechanical activity was blocked with TTX (Fig. 1B and C). Overall, between days 5 and 9, total AChE rose 11-fold in fibrillating and only 2-fold in nonfibrillating myotubes (Fig. 1D).

To evaluate the effect of spontaneous electromechanical activity on individual globular and asymmetric forms, the globular and asymmetric fractions obtained by sequential extraction were applied to sucrose gradients. The gradient profiles obtained from one set of myotube cultures are shown in Figures 2 (5-day cultures) and 3 (8-day cultures). At 5 days, the activities of the 10 S and 4 S forms were the same in control myotubes (which were just beginning to fibrillate) and in myotubes treated with TTX (Fig. 2, A and C). Both globular forms increased in fibrillating and nonfibrillating myotubes between 5 and 8 days, but the increases in the fibrillating myotubes were greater. By 8 days, fibrillating myotubes contained 4.8-fold more 10 S (G1) and 4.3-fold more 4 S (G2) AChE (Fig. 3, A and C) than did nonfibrillating myotubes, with total levels of globular forms being 4.4-fold greater in fibrillating myotubes. At 5 days, the asymmetric fraction was composed of three forms that sedimented at 16 S and at approximately 10 S and 6 S. The activity of each of these forms was essentially identical in fibrillating and nonfibrillating myotubes at this stage (Fig. 2, B and D). The asymmetric forms in fibrillating myotubes increased dramatically between 5 and 8 days, but there was no appreciable change in any of the asymmetric forms in nonfibrillating myotubes. By 8 days, fibrillating myotubes had 19.3-fold more 16 S (A12) AChE, they contained appreciable 12.5 S (A3) AChE that was not apparent in the nonfibrillating cells, and they contained increased amounts of the 10 S form found in the asymmetric fraction (Fig. 3, B and D). At 8 days the fibrillating myotubes had 11.2-fold more total asymmetric AChE than did nonfibrillating myotubes. The 6 S form in the asymmetric fraction was difficult to evaluate in fibrillating myotubes, but it clearly did not increase as the other asymmetric forms did.

![Figure 1. Development of AChE forms in myotube cultures. Myotube cultures at various ages were subjected to sequential extraction. AChE activity then was measured. Globular, asymmetric, nonextractable, and total AChE are shown in A, B, C, and D, respectively. Data for untreated myotube cultures (O), myotube cultures treated at day 4 with 1 µM TTX (O), TTX-treated myotube cultures washed at day 7 to remove TTX (A), and myotube cultures to which TTX was added at day 7 (Δ) are shown. Activity is expressed as picomoles of ACh hydrolyzed/min-' dish-' . Each point represents the mean ± SEM of three to eight determinations. Where no error is shown the error is less than the size of the symbol.](attachment://figure1.png)
Morphological development. The effect of TTX on AChE was not related to changes in the gross morphological development of the myotubes. The percentage of total surface area of the culture dish covered by myotubes was carefully examined in 5- to 8-day-old cultures as described under "Materials and Methods." The surface area covered by myotubes increased linearly from 13% to 35% in both fibrillating and nonfibrillating cultures (Fig. 4).

Reversibility of TTX effect. The effect of TTX was reversible. When cultures that had been exposed to TTX on day 4 were washed on day 7 and bathed in medium obtained from 7-day fibrillating cultures (conditioned medium), globular forms returned to control levels in 2 days (Fig. 1A). Asymmetric (Fig. 1B) and non-extractable (Fig. 1C) forms also recovered dramatically over 2 days. We used conditioned medium rather than fresh medium in this experiment because AChE activity decreases when medium that has been conditioned for several days is replaced with fresh medium (Table I). Control experiments established that the increase in AChE activity was due to the resumption of electromechanical activity rather than to factors in the conditioned medium. When medium from fibrillating myotube cultures was supplemented with TTX and used to replace the medium of the nonfibrillating cultures, AChE activity did not increase (Table I).

Inhibition of fibrillation at 7 days (when cultures were fibrillating vigorously and had developed significant levels of AChE) not only prevented AChE from increasing normally but also caused all fractions to be reduced
within 1 day to levels only slightly higher than those found in cultures where fibrillation had been entirely prevented (Fig. 1). Thus the effect of TTX is reversible, occurs relatively rapidly, and is not critically dependent on the developmental stage of the culture.

**Titration of AChE active sites with MTP.** The increase in AChE activity associated with fibrillation could be due either to an increase in the activity of individual AChE molecules or to an increase in the number of molecules present. Vigny et al. (1978) have shown that the stable, irreversible phosphorylating agent MTP can be used to titrate AChE active sites stoichiometrically. MTP binds to AChE active site extremely rapidly (the second-order rate constant is about $1.4 \times 10^6 \text{ mol}^{-1} \text{ set}^{-1} \text{sec}^{-1}$) and completely irreversibly with each molecule of inhibitor added to the homogenate inactivating one active site. We used MTP to evaluate the number and catalytic activity in AChE active sites in 8-day fibrillating and nonfibrillating myotubes. In these experiments, homogenates of fibrillating and nonfibrillating myotubes having the same AChE activity were prepared as described under "Materials and Methods." An identical amount of MTP (estimated to inactivate half of the active sites) then was added to the two homogenates, and they were assayed periodically for residual AChE activity. Typical results were shown in Figure 5. It is evident that MTP had an essentially identical effect on the AChE from fibrillating and nonfibrillating myotubes. This result provides strong evidence that the catalytic activity of individual active sites is essentially identical in fibrillating and nonfibrillating myotubes, so that the increase in AChE activity associated with fibrillation is due primarily, if not exclusively, to an increase in the number of AChE molecules present. In three experiments on 8-day cultures, each dish of fibrillating myotubes contained 0.53 ± 0.08 pmol of active sites and each dish of nonfibrillating myotubes contained 0.10 ± 0.02 pmol of active sites. The apparent first order rate constant for acetylcholine turnover at active sites in fibrillating myotubes ($229 \pm 22 \text{ sec}^{-1}$) was not different statistically from that in nonfibrillating myotubes ($217 \pm 9 \text{ sec}^{-1}$). This turnover number for rat myotube AChE corresponds to a value that is one-third to one-half of that reported by Vigny et al. (1978) for rat brain AChE after allowance is made for the different assay conditions used in our study.

**Metabolism of AChE.** The increased number of AChE molecules in fibrillating myotubes could be due to increased synthesis, decreased turnover, or to some combination of the two. To evaluate AChE metabolisms, we examined the enzyme after inactivating it with paraoxon (diethyl p-nitrophenyl phosphate). Paraoxon readily penetrates cells and diethylphosphorylates a serine in the active site of AChE, thereby inactivating the enzyme and releasing p-nitrophenol. The diethylphosphorylated enzyme is stable, but it can be reactivated rapidly and quantitatively with 2-PAM (Brockman et al., 1982; Younkin et al., 1982). Thus one can inactivate AChE with paraoxon and simultaneously evaluate synthesis (by following the newly synthesized active AChE) and turnover (by following the 2-PAM-reactivatable AChE). The combination of diethylphosphorylation and 2-PAM reactivation provides, in effect, an extremely sensitive and highly specific way to label AChE. An attractive feature of this method is that there is no reutilization of label following degradation of labeled AChE or following spontaneous reactivation of inactivated (labeled) AChE because the enzyme is not inactivated (labeled) by diethylphosphate per se.

In homogenates in vitro, the rate of spontaneous reactivation of diethylphosphorylated AChE is about 0.026 hr\(^{-1}\) at 37°C (Newman et al., 1983) Therefore, there should be very little generation of active AChE by reactivation of the labeled pool in short experiments like those described below. To confirm this, control experiments were performed in which cellular metabolism was inhibited as described under "Materials and Methods." The active enzyme that appeared in these cultures after removal of paraoxon represents the maximal amount that could be generated by spontaneous reactivation of diethylphosphorylated enzyme. Over a 2-hr period, only
104 pmol min$^{-1}$ dish$^{-1}$ of active enzyme appeared in the cultures ($n = 4$). This represented only 5.1% of the total amount of labeled enzyme present immediately after treatment with paraoxon. Thus the spontaneous reactivation of diethylphosphorylated AChE in cultured myotubes occurs slowly and at essentially the rate observed in vitro.

Control experiments were performed to evaluate the impact of paraoxon on AChE metabolism. If paraoxon inactivation influenced AChE metabolism appreciably, then the amount of globular and/or asymmetric AChE present would change. Labeling by paraoxon inactivation is advantageous because the total amount of AChE present can be assessed by assaying AChE after reactivating the diethylphosphorylated enzyme with 2-PAM. Figure 6 shows that no appreciable change in total globular or asymmetric AChE (measured after 2-PAM reactivation) occurred in either fibrillating or nonfibrillating myotubes over the 2-hr period of our experiments.

Seven-day fibrillating and nonfibrillating cultures were treated with 10 $\mu$M paraoxon for 30 min to label (diethylphosphorylate) AChE. This protocol labeled an average of 86% of the enzyme in the cells. The paraoxon was then washed from the cultures and, at various times, the myotubes were harvested. The cells were homogenized and two fractions were prepared. One contained globular AChE and the other contained enzyme (left after extraction of the globular enzyme) that is almost entirely asymmetric AChE. These fractions were assayed before and after reactivation of the diethylphosphorylated enzyme with 2-PAM in order to evaluate both newly synthesized (active) AChE and labeled (2-PAM-reactivable) enzyme.

The percentage of labeled globular and asymmetric AChE remaining at various times after labeling is shown in Figure 7. The half-life of globular AChE was shorter in fibrillating cells (85 min) than in nonfibrillating cells (122 min), but the difference was not statistically significant. The turnover rate of asymmetric AChE was too slow to assess accurately in the 2-hr interval over which our measurements were made. In Figure 7 it appears that asymmetric AChE turns over more slowly in fibrillating than in nonfibrillating myotubes. The difference between fibrillating and nonfibrillating cells was not significant, however, and, in a second series of experiments in which turnover rates were approximately the same as those in Figure 7 (data not shown), asymmetric AChE turned over more rapidly in fibrillating than in nonfibrillating myotubes (again the difference was not significant).

The newly synthesized AChE measured at various times after inactivation is shown in Figure 8. It is apparent that newly synthesized globular and asymmetric forms both appear more rapidly in fibrillating than in nonfibrillating myotubes. Based on the reappearance of AChE in the first 60 min, it appears that globular forms are synthesized 7 times faster in fibrillating myotubes than in nonfibrillating myotubes and that asymmetric forms are synthesized 6 times faster in fibrillating than in nonfibrillating myotubes.

The "asymmetric" AChE evaluated above was the enzyme left after a single extraction of globular forms. This enzyme consists mostly of soluble asymmetric forms but also contains nonextractable AChE and some residual globular forms. To rule out the possibility that increased synthesis was occurring entirely in the contaminating forms, we examined the rate of synthesis of the soluble asymmetric AChE obtained after extracting twice
to remove globular forms (see "Materials and Methods" for details of the extraction procedure). Figure 9 clearly shows that soluble asymmetric AChE is synthesized more rapidly in fibrillating than in nonfibrillating myotubes. Following inhibition, asymmetric AChE appeared about 4 times faster in fibrillating than in nonfibrillating myotubes over the first 2 hr of recovery.

**Discussion**

On the basis of evidence obtained in earlier work, we proposed (Collins and Younkin, 1982) that the electromechanical activity set up in mammalian skeletal muscle by motor neurons increases nonjunctional AChE by increasing the synthesis of the globular forms of the enzyme. We suggested, in addition, that increased synthesis of asymmetric forms might contribute to the effect of motor neurons on junctional AChE. In this study, we have tested the hypothesis that electromechanical activity increases the synthesis of globular and asymmetric AChE by comparing fibrillating rat myotubes with TTX-treated nonfibrillating myotubes.

Our experiments confirm many of the results reported by Rieger et al. (1980) in their study of the effect of fibrillation on cultured embryonic rat myotubes. Like Rieger et al. (1980) we find (1) that a large increase occurs in the 16 S (A12) form of AChE (19-fold in our study) between 5 and 8 days when fibrillation occurs in the myotubes, (2) that this increase in A12 AChE is prevented or reversed by preventing or stopping fibrillation with TTX, and (3) that the effect of TTX is reversible (Figs. 1 to 3).

In their study, Rieger et al. (1980) concentrated on the A12 form of AChE and did not report on the other forms of the enzyme. Our data indicate that the 4 S (G1), 10 S (G4), and 12.5 S (A6) forms are also affected when fibrillation is blocked with TTX. Between 5 and 8 days in culture, fibrillating myotubes develop over 4 times more 4 S (G1) and 10 S (A6) AChE than do nonfibrillating myotubes, and the 12.5 S (A6) form, which is not detectable in nonfibrillating myotubes, develops appreciably in fibrillating cells between 5 and 8 days (Figs. 2 and 3). Thus, inhibition of fibrillation with TTX substantially impairs the development of both globular (G4 and G1) and asymmetric (A12 and A6) forms of AChE with a much larger and more dramatic effect being exerted on the asymmetric forms.

Our results differ from those reported by Rieger et al. (1980) in one relatively minor way. In their study, no asymmetric forms were detected when fibrillation was prevented. In the asymmetric fraction from our nonfibrillating myotubes (Figs. 2D and 3D), three distinct forms were clearly present: the classical 16 S (A12) form and two other forms that sedimented at about 10 S and 6 S. Gradient fractions were not assayed in the presence of iso-OMPA, so that it is possible that the two smaller forms in the asymmetric fraction are not true AChE. It appears that the 10 S form in the asymmetric fraction is not simply residual globular enzyme because it, like the 16 S (A12) and 12.5 S (A6) forms, aggregates and migrates to the bottom of low ionic strength sucrose gradients whereas the 10 S form in the globular fraction does not (Brockman et al., 1982). We have not yet investigated the 6 S form in the asymmetric fraction in this manner. In our experimentation, we isolated the asymmetric fraction before separating the individual asymmetric forms by velocity sedimentation. Rieger et al. (1980) separated both globular and asymmetric forms on the same gradient. This makes it impossible to detect the unusual 10 S and 6 S forms and difficult to detect the classical 16 S form. Figure 2, C and D, shows that the 16 S form in nonfibrillating myotubes is a tiny fraction of the 10 S and 4 S forms. It seems quite possible that this form was lost in the base line when it was examined on the same gradient as the more abundant globular forms. Our study...
was carried out on myotubes derived from 20-day embryos whereas Rieger et al. (1980) examined myotubes derived from 18-day embryos. This may also account for the minor difference in the two studies.

The AChE level in 7-day fibrillating myotubes decreases when the culture medium is replaced, presumably because of the removal of factor(s) released into the medium by fibrillating cells. This raised the possibility that the effect of electromechanical activity is mediated by factor(s) released into the medium. Table I shows, however, that conditioned medium by itself had little effect on AChE levels in nonfibrillating myotubes. Thus electromechanical activity and conditioned medium are both required for the development of maximal AChE activity.

Our morphological data (Fig. 4) show that the effect of TTX is not due to impairment of the gross morphological development of the myotubes. The experiments in which we used MTP to titrate AChE active sites (Fig. 5) establish that the increased AChE activity caused by fibrillation is due to an increase in the number of AChE molecules present and not to an increase in the rate at which individual AChE molecules turn over acetylcholine.

Our examination of AChE metabolism shows that both globular and asymmetric forms of AChE are synthesized more rapidly in fibrillating than in TTX-treated nonfibrillating myotubes. Newly synthesized globular forms of AChE accumulated much more rapidly in fibrillating than in nonfibrillating myotubes (Fig. 8), and the turnover of these forms was, if anything, more rapid in fibrillating than in nonfibrillating cells (Fig. 7); thus the increased level of globular AChE in fibrillating myotubes appears to be due entirely to increased synthesis. The asymmetric forms turned over very little in the 2-hr interval over which metabolism was measured; therefore the turnover of this pool could not be accurately measured. The increased accumulation of newly synthesized asymmetric forms that was observed in fibrillating myotubes (Figs. 8 and 9) clearly indicates that increased synthesis contributes to the increased level of asymmetric forms in fibrillating myotubes.

There have been two studies showing that electromechanical activity decreases the synthesis of acetylcholine receptors in mammalian skeletal muscle (Reiness and Hall, 1977; Linden and Fambrough, 1979). Our data provide strong evidence that electromechanical activity increases the synthesis of globular and asymmetric forms of AChE. It appears, therefore, that electromechanical activity can regulate the synthesis of the functionally important surface proteins in skeletal muscle in either a positive or a negative direction.

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