MYOSIN LIGHT CHAIN 1 ISOFORM EXPRESSION REMAINS CONSTANT DURING AGEING IN WISTAR F455 RATS

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(Received January 3rd, 1991)
(Revision received April 23rd, 1991)

SUMMARY

In order to study muscle gene expression during ageing, we examined both protein and total cellular RNA from Wistar F455 rat soleus and extensor digitorum longus (EDL) muscles at a variety of chronological ages. We found no evidence of the reappearance of the fast protein isoform of myosin light chain 1 [MLC1] in the slow soleus muscle during ageing previously reported by Syrovy and Gutmann, Pflügers Arch., 369 (1977) 85-89. We used both SDS-PAGE analysis of MLC1 proteins and slot blot RNA analysis with a probe specific for rat fast MLC1 mRNA (pC91), and found no changes in fast MLC1 expression during ageing in soleus or EDL muscles from these rats. These results indicate that re-expression of the fast MLC1 isoform is not a universal property of ageing soleus muscle.

Key words: Myosin light chain 1; Rat; Ageing; RNA; SDS-PAGE; Muscle isoform(s)

INTRODUCTION

Mammalian skeletal muscle alters histochemically and morphologically during chronological ageing [1-8]. The causes and regulation of these age-related changes are not well understood. To address the role of the body environment in the muscle ageing process, Carlson and Faulkner [9] used a cross-age muscle grafting system. This study examined morphological and physiological changes in soleus and exten-
sor digitorum longus (EDL) muscles transplanted between young and old animals. The authors reported that the age of the host animal is more important in determining the apparent morphological and physiological ‘age’ of the grafted muscle than the chronological age of the muscle itself, supporting the idea that the environment somehow affects muscle ageing.

The events leading to the changes which constitute ‘ageing’ have not been elucidated. No qualitative or marked quantitative change in gene expression linked to the ageing process has yet been identified [10]. Any such molecular change would be of real advantage in assessing the effect of epigenetic factors on muscle ageing. It could potentially serve as an ‘ageing indicator’ to test whether, for example, gene expression alters in cross-age muscle transplants, resulting in morphological and physiological changes dictated by their new environment. If muscle protein or gene expression changes qualitatively with age, as opposed to quantitative changes due to changes in mRNA stability or protein synthesis rates, the factors which control such changes during ageing could be studied, with a view to understanding the ageing process as a whole.

In 1977, Syrovy and Gutmann [8] reported that during ageing, protein of the fast isoform of MLC1 (myosin light chain 1) began to reappear in rat soleus muscles. This isoform, produced by an alternative splicing event [11], normally appears perinatally in both rat soleus and EDL muscles but disappears in the soleus as this slow muscle matures. If the gene for the fast isoform of MLC1 were re-expressed in aged soleus, such an age-related molecular change would be particularly useful as an ageing marker due to the high expression level of myosin in skeletal muscle [12] and the availability of several techniques to detect distinct isoforms of this well-studied protein [12,13], ranging from SDS-PAGE and antibody detection to Northern analysis with cDNA probes [11,14–16].

Given that Syrovy and Gutmann’s report [8] described a promising candidate for an ageing marker, we examined aged soleus muscles for this reported reappearance of fast MLC1 protein. In the present study, we found no evidence for any alteration in MLC1 isoform expression in soleus muscle during ageing, and therefore we conclude that, even if this change reproducibly occurs in the strain of rats studied by Syrovy and Gutmann, it is not a universal age-related phenomenon.

MATERIALS AND METHODS

Dissection

Thirty-nine Wistar F455 rats kindly donated by Dr. Bruce Carlson from his colony at the University of Michigan were killed by CO₂ inhalation; the soleus and EDL muscles were immediately removed from both hindlimbs, minced, and placed in sterile preweighed tubes. The tubes were reweighed, frozen in liquid nitrogen, and stored at -85°C until use.
**SDS-PAGE sample preparation**

The protocol followed for SDS-PAGE was a modification of Laemmli [17] developed by Esser et al. [18] and communicated to us by J. Opiteck of the Division of Kinesiology at the University of Michigan; the solutions used are according to their protocol. Frozen minced muscle was homogenized in 1 ml of homogenizing buffer (62.5 mM Tris, pH 8.0; Sigma) per 100 mg muscle using a Tekmar homogenizer. This 100 mg/ml muscle homogenate stock was stored frozen at −85°C.

To determine protein content, an aliquot of this stock was incubated at 65°C overnight (12–20 h) in 0.25 N NaOH to hydrolyze protein and prevent aggregation without inhibiting amino acid-Coomassie blue binding during the Bradford protein assay. Each aliquot was neutralized with 0.25 N HCl. A standard protein assay [19] was performed using Bio-Rad's protein assay reagent and bovine serum albumin (fraction V; Sigma) as the standard. Averaged values of triplicate readings were used to determine protein concentration in each stock sample.

Another aliquot of each muscle homogenate stock was combined 1:1 (v/v) with 2× sample buffer [18], boiled for 3–5 min, cooled on ice, and frozen at −85°C until diluted for use on polyacrylamide gels. Gel samples were diluted to a protein concentration of from 0.67 to 2.67 mg/ml with 1× sample buffer. These samples were stored either at −20°C (2 weeks or less) or at −85°C.

**SDS-polyacrylamide gels**

We adapted the Esser et al. [18] protocol for use with miniature gradient gels; the small samples required for these gels allowed us to analyze samples from individual muscles repeatedly. Ten to twenty percent gradient gels (80 × 50 × 0.75 mm) were cast using a BioRad MiniProtean II gel apparatus and a Buchler Instruments conical-chamber gradient former. A five percent stacking gel was used. The upper buffer contained 25 mM Tris, 192.5 mM glycine, 0.1% SDS; pH 8.3. The lower buffer contained 375 mM Tris, 0.1% SDS; pH 8.8. The upper buffer was cooled and electrophoresis carried out at 14°C. The gels were run at 150 V for 120 min and stored in 40% methanol/10% acetic acid until staining.

**Gel staining**

The Bio-Rad silver stain solutions and protocol were followed, with suggested adjustments to increase sensitivity and decrease background. The gels were stored in distilled, deionized water.

**Gel photography**

A Kodak Electrophoresis Duplicating Film Kit was used to produce clear positives of selected gels. Optimum exposure times ranged from 35 to 45 s, using a Bausch and Lomb 8 W 115 V 60 cycle fluorescent lamp at 8 inches distance from the gel.
**Film scanning**

Each film was scanned in a Gilford Response spectrophotometer. The parameters used to scan the films were held constant (bandwidth 0.5 nm; wavelength 500 nm; slitplate 0.05 × 2.36 mm). The clear area of the film at the gel’s 20% end was used to establish background readings in each lane, and each lane was then scanned up the gradient, from the 20% to the 10% region.

**Slot blot analysis of RNA**

After confirming with SDS-PAGE analysis that the MLC1 band patterns of male and female muscles were alike, total cellular RNA was isolated from male and female Wistar F455 rats’ soleus and EDL muscles using the RNAzol solution and protocol (Cinna-Biotecx). These RNA samples were then blotted onto Nytran membrane using a slot blottter (Schleicher and Schuell) [21]. RNA samples isolated from myotubes of the C2C12 cell line (developed by Yaffe and Saxel [22]; subcloned by Blau et al. [23]), rat gastrocnemius (a mixed fast- and slow-twitch muscle), rat heart, and rat brain were used as controls.

Restriction fragments from plasmid digests were random-primer labeled with [32P]dCTP (ICN) by the method of Feinberg and Vogelstein [24,25] for use as probes. The fragments used were the PstI/EcoRI fragment of pC91-pUC18Bam− (courtesy of N. Rosenthal), a rat cDNA fragment specific to the fast MLC1 [11]; the PstI fragment of pCl10.4 (courtesy of M. Buckingham), a mouse cDNA fragment specific to the embryonic isoform of MLC1 [MLCemb] [26]; and the EcoRI/HindIII fragment of pPG26 (courtesy of R. Wade), a BglII/KpnI fragment of the human skeletal muscle cDNA homologous to *Drosophila* heat shock protein 83 [HSP83] [27].

Blots were probed as described in Sambrook et al. [21], but hybridization was performed at 42°C (pCl10.4 and pPG26) or 50°C (pC91) for 10–20 h in 50% formamide, 5× Denhardt’s solution, 0.1% SDS, ~200 mg/L sheared salmon sperm DNA, and 5× SSPE. After probing with the MLCemb probe (pCl10.4), blots were washed with increasing stringency, the last wash containing 0.1× SSC and 1% SDS at 50°C. Blots were exposed to Kodak XAR-5 X-ray film for 50 h with DuPont Cronex intensifying screens at ~85°C, and developed in a Kodak X-omat automatic film processor. Blots were then stripped in 5 mM Tris, pH 8.0; 0.2 mM EDTA; 0.05% pyrophosphate; 0.1× Denhardt’s solution and then reprobed with the fast MLC1 probe (pC91), as above, and washed to a final stringency of 0.1× SSC and 1% SDS at 67°C. They were exposed to film for 44 h and developed as above. After stripping, the blots were reprobed once again with the HSP83 probe (pPG26), washed to a final stringency of 1× SSC and 1% SDS at 50°C, and exposed to film for 3 days.

**RESULTS**

**Protein analysis**

Our modification of the Esser et al. [18] protocol for SDS-PAGE employs in-
dividually poured gradient minigels. Such an analysis allows precise determination of the MLC1 isoform expression in small quantities (5–100 µg) of total muscle homogenate from individual muscles. Each 15-lane minigel is 80 × 50 × 0.75 mm; its small size and steep gradient (10–20%) create sharp, well-resolved, scannable bands while reducing the amount of muscle used per lane. The use of total muscle homogenate (without the extensive purification measures used by others [8]) eliminates the possibility that isoform ratios will be altered by selective precipitation of myosin isoforms during the purification process.

Muscle samples run on each gel included in most cases at least one example of young soleus, young EDL, old soleus, and old EDL, in order to ensure clear identification of specific MLC bands (Fig. 1). Within each gel, the lane-to-lane variation in migration distances was extremely small, and the use of gradient gels produced very highly resolved bands while retaining a wide weight range of proteins on the gel. Due to the reproducibility of the characteristic band patterns of the control (young) soleus and EDL samples included on each gel, gel-to-gel comparison was possible.

Spectrophotometric scans confirmed the reproducibility and good resolution of the gels and allowed the determination of standard migration distances for the fast and slow isoforms of MLC1 on each gel by comparison of the band patterns with published data [28, K.A. Esser, pers. commun.]. The scan data were examined for appearance of the fast MLC1 isoform in either young adult or old soleus; in contrast to Syrovy and Gutmann’s report [8], no fast MLC1 band peaks were found in any soleus sample (Fig. 2). A second set of gels was run using samples from female rats to establish whether any gender-specific difference exists in the soleus muscle’s response to ageing. No differences in MLC1 expression were noted in these gels (data not shown).

![Fig. 1. Typical polyacrylamide gradient minigel showing the fast and slow isoforms of MLC1. Mf = fast isoform, Ms = slow isoform. Each sample is loaded in three different concentrations (0.67, 1.33, 2.67 µg/lane: referred to as 1 x, 2 x, 4 x). L = molecular weight ladder; OS = old soleus; YE = young soleus; YE = young EDL standard. Starred lanes correspond to the scans shown in Fig. 2.](image-url)
Fig. 2. Scans (taken of the lanes starred in the above gel) showing expected locations of fast and slow isoforms of MLC1, as determined by lane-to-lane and gel-to-gel comparisons. $M_R$ = reference peak, used as a location reference to normalize migration distances within each gel; $M_f$ = the fast isoform's expected location; $M_s$ = the slow isoform's expected location. A: young soleus; B: young EDL; C: old soleus. Note the absence of any fast MLC1 peak in old soleus.
Fig. 3. Three sequential probings of a typical slot blot containing RNA from male and female rats aged 10 to 107 weeks. Each slot contains 2 µg total cellular RNA. Slots marked ‘- - -’ contain no RNA. A: Results of the fast MLC1 probe. Note strong binding to EDL and gastrocnemius (mixed fast/slow muscle) only, with no binding to soleus, C2C12 myotube, heart, liver, or brain. B: Results of the embryonic MLC probe. Note that this probe is specific to C2C12 myotube RNA only, with no cross-reactivity to any adult rat MLC’s. C: Results of the human homologue to the Drosophila HSP83. Note similar expression levels in all samples, in contrast to the other two probes.
RNA analysis

We confirmed the results of our SDS-PAGE analysis by analyzing total soleus and EDL muscle RNA on slot blots, using probes specific to fast MLC1, embryonic MLC1, and human HSP83 as a control for loading variations (Fig. 3). RNA isolated from male and female rats of a variety of ages ranging from 10 to 107 weeks was analyzed using slot blotting techniques with cDNA probes specific for MLC isoform mRNA. The MLC_{emb} isoform was found to be expressed only in the C2C12 myotube control; no hybridization was seen in any of the adult rat muscle RNA samples which we examined. The fast MLC1 isoform was expressed exclusively in EDL muscles of all ages examined, with no mRNA detected in soleus samples of any age. It was also present in rat gastrocnemius, a mixed fast and slow fiber muscle, but was not found in rat heart or brain. Consistent loading was confirmed by probing with a cDNA probe to the human heat shock protein 83, which bound comparably to all samples and was possibly more evident in soleus than in EDL samples, in contrast to the MLC1 fast probe. This result ensured that sufficient RNA was present to have detected any expression of fast MLC1 in soleus.

DISCUSSION

Our data show that in Wistar rats from the University of Michigan colony, fast MLC1 expression remains very low or nonexistent in both male and female rat soleus muscles during ageing, and no embryonic isoforms are found at any of the adult ages examined. This result is in marked contrast to Syrovy and Gutmann's report [8], which shows a distinct increase in fast MLC1 protein in aged rat soleus muscle. The differences in our data may result from variations in rat strains; however, their paper did not report the strain of rat used in their experiments. Because our local colony of rats has been inbred for over 90 generations and thus may vary from the standard Wistar strain [29], strain differences remain one possible explanation.

Another possible cause of the discrepancy between our results and Syrovy and Gutmann's may be that our gel method used unpurified myosin samples, whereas Syrovy and Gutmann used a standard purification protocol to separate myosins from other muscle proteins before applying their samples to the gel. This purification procedure may alter the relative amounts of slow and fast MLC1 isoform present in the gel sample. Because our RNA analysis showed no trace of fast MLC1 in old soleus, confirming our protein analysis, we believe that our unpurified samples yielded accurate results.

Because we do not have access to the strain of rats used by Syrovy and Gutmann, nor to a complete description of their rat care conditions, we cannot determine whether strain or care differences are the possible cause of these conflicting data. Rat strain differences have been reported in other studies of rat skeletal muscle [5]. While no reports of rat care inducing a specific genetic effect, such as the reappearance of
the fast MLC1 isoform, are available, care conditions have been shown to correlate with age-related effects [30]. Thus, although our rats were not raised in a specific pathogen-free, barrier-protected environment, differences in either rat care or rat strain might account for the differences between our results and Syrovy and Gutmann’s.

Because the ‘dedifferentiation’ (the age-related reappearance of a MLC1 isoform normally absent in soleus after maturation) noted by Syrovy and Gutmann does not appear to be a universal phenomenon, the re-expression of fast MLC1 isoform in rat soleus muscle cannot be used as a molecular ‘age marker’ in studies of muscle ageing. Furthermore, our study raises the possibility that changes in gene expression at the transcriptional level, at least in the case of MLC1 isoforms, may not be the normal underlying cause of histochemical and morphological changes in rat skeletal muscle during ageing. While many studies have shown changes in muscle histochemistry and morphology during ageing [1-8], alterations in gene expression which can directly account for these changes have not been demonstrated. Epigenetic effects may thus turn out to play a much larger role in the ageing process than previously supposed [10].

ACKNOWLEDGEMENTS

We would like to thank B. Carlson for donating rats for this study, J. Opiteck and K. Esser for SDS-PAGE instruction and advice and A.I. Kavka and C.A. Gardner for helpful comments on the manuscript. This work was supported by grants from the NIH, the NSF, the MDA and the Rackham Foundation of the University of Michigan to KFB; grant #GM30866 from the NIH to MIL; and grants from the Cancer Research Committee and the Biomedical Research Council of the University of Michigan to KFB and MIL.

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