Incorporation of Nascent Myosin Heavy Chains into Thick Filaments of Cardiac Myocytes in Thyroid-treated Rabbits

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Abstract. A monoclonal antibody (mAb 37) specific for α-myosin heavy chain (α-MHC) is used to follow the spatial and temporal incorporation of α-MHC into rabbit left ventricular myocytes. The expression of the two adult cardiac MHC genes, α and β, is regulated by manipulating the thyroid hormone level of the animal. 10 wk on a propylthiouracil diet down-regulates expression of α-MHC to near 0%. α-MHC gene expression is up-regulated by injecting L-triiodothyronine (100 μg/kg per d) for 1-4 d. This protocol provides a means by which to follow the redistribution pattern of α-MHC within the myocyte in vivo. A uniform distribution of immunofluorescent signal is seen within every myocyte throughout the left ventricle. Ultracryomicrotomy without fixation is used to obtain sections for immunogold-electron microscopy. To quantify the immunogold method the density of gold-labeled antibody per unit of area tissue is determined for various regions of the sarcomere. Tissue from normal and 2-wk baby has a uniform distribution of gold density along the length of the A band. The average gold density of the A band increases with days of thyroid injection from 38 ± 4 grains/μm² (n = 2 animals) (mean ± SE) at day 1 to 182 ± 59 grains (n = 2 animals) at day 4. There is a nonuniform incorporation of the newly synthesized α-MHC within the A band of thyroid-treated animals since 50% more of the α-MHC is found at the ends of the A band while the center of the A band has 40% less than the average α-MHC content (grains/μm², n = 7 animals). These results support a thick filament assembly model that allows every myosin in a thick filament to be exchanged with new myosin. However, in the intact functioning myocyte, there is greater exchange of new myosin at the ends than in the central region of the thick filament.

The thick filament of muscle is composed of approximately 300 myosin molecules whose function is to generate force during muscle contraction. Cardiac myosin is degraded and replaced with a half life of 8 d in the rabbit (Morkin et al., 1972) and the turnover process is accomplished without interrupting the force-generating function of the muscle fiber. Because of this special constraint, the nature of thick filament assembly and the mechanism involved with exchange of myosin within that filament have long been areas of scientific inquiry.

Huxley (1963) showed that free myosin molecules could spontaneously form thick filaments when the ionic strength of the solution was lowered. Further work on the self-assembly process revealed that strong ionic interactions exist between the rod regions of myosin molecules and are responsible for spontaneous assembly (for review see Harrington and Rodgers, 1984). In an attempt to understand the mechanisms that may regulate thick filament assembly, numerous investigators have studied the system in vitro by varying pH, ionic strength (for review see Pepe, 1983; Harrington and Rodgers, 1984), temperature, calcium concentration (Higuchi and Ishiwata, 1985), and in the presence of sodium pyrophosphate (Reisler et al., 1986). Saad et al. (1986a, c) studied the rapid kinetics of thick filament assembly with the fluorescence energy transfer system and found that a monomeric pool of myosin is in a state of dynamic exchange with thick filaments. However, synthetic thick filaments lack many of the structural and functional elements of the living muscle fiber that might alter both the kinetics of the assembly process and the size of the free myosin pool.

One reason why myosin exchange has not been studied in vivo is the difficulty in localizing newly synthesized myosin within the muscle fiber. In the past the only method available was by autoradiographic analysis of tissue that had been infused with radioactive amino acid precursors. Unfortunately, this method labels all new proteins nonspecifically and can not identify the cellular distribution of myosin alone.

The design of the present experiment takes advantage of the fact that cardiac myosin heavy chain (MHC) exists in two isoforms, α and β, and the expression of the isoforms can be regulated by manipulating the thyroid level of the animal. Investigations have shown that the level of α-MHC expression can be completely down-regulated by inducing a state of hypothyroidism in the experimental animal (Hoh et al., 1978). We found that 10 wk at a low plasma thyroid level...
(100 ng/dl) was needed to switch off the α-MHC gene for a sufficient period of time to ensure that all preexisting α-MHC protein was removed. Up-regulation of α-MHC gene is accomplished by increasing the plasma thyroid level (Chizzonite et al., 1982) with injections of L-triiodothyronine.

A second aspect that is critical to the design of this study is the availability of a method to differentiate between the α and β isoforms of MHC. Even though the isoforms share 95% homology of amino acid sequence (Umeda et al., 1986), a monoclonal antibody, mAb 37, specific for the α-MHC isoform has been developed (Chizzonite et al., 1982). The sites of incorporation of α-MHC into the thick filament are monitored by immunogold-electron microscopy.

There are many possible patterns of incorporation of new myosin into thick filaments. One pattern would be de novo synthesis of new filaments to replace old. Another is a random incorporation of new myosin along the length of the filament. A third is preferential addition of new myosin at some site along the thick filament. Our results show that during the period of induced isomyosin switching nascent myosin is incorporated throughout the length of the thick filament. However, a preferential addition of the nascent myosin to the ends of the thick filament is also observed.

**Materials and Methods**

**Thyroid Model and Tissue Preparation**

10 New Zealand White rabbits (2.0–3.0 kg) were placed on a propylthiouracil (PTU)-enriched diet (Teklad Diets, Madison, WI) for 70 d to induce hypothyroidism (Everett et al., 1984). At the end of this time the rabbits were injected intraperitoneally with L-triiodothyronine (l-T3; Sigma Chemical Co., St. Louis, MO) in saline solution (1 mg/ml) at a dose level of 100 μg/kg per d for 1–4 d. The animals were killed by the infusion of a high potassium solution (Kovacs et al., 1983) into the vena cava to depolarize the heart and produce a state of relaxation. The heart was removed, the ventricles dissected, blotted dry, and weighed. The papillary muscles of the left ventricle were dissected free from the heart wall and pinned down at rest length to prevent contraction during subsequent dissection and sucrose infiltration.

**Immunofluorescence**

Procedures for indirect immunofluorescence were as presented in previous work (Eisenberg et al., 1985). Frozen sections were incubated in mAb 37 diluted 1:100 (a gift from Dr. R. Zak, University of Chicago, Chicago, IL), and in goat anti-rat IgG FITC (bottom) 5 nm gold spheres (Janssen Life Instruments, Inc., Gaithersburg, MD) at -110°C with a tungsten-coated glass knife (Roberts, 1975). Sections were retrieved with a drop of 2.3 M sucrose containing 0.75% gelatin (type III, Sigma Chemical Co.). The gelatin reduced the surface tension of the sucrose drop as it thawed, which decreased but did not eliminate fragmentation of the unfixed tissue.

**Ultracytomicrotomy**

The methods of Tokuyasu (1986) were followed for both ultracytomicrotomy and immunostaining. Small blocks of tissue from the papillary muscles were immediately cryoprotected with 2.3 M sucrose in 0.1 M PBS. The antibody, mAb 37, no longer bound to the antigeneic site after the following fixations: 0.2% glutaraldehyde (perfused or immersed); 3% paraformaldehyde; paraformaldehyde/lysine/sodium m-periodate (PLP fixation) (McLean and Nakane, 1974); 1.5% glutaraldehyde and 20 mM ethylendiaminetetraacetate (Tokuyasu et al., 1981); 0.3% glutaraldehyde, 1.4% Tris, 50% PBS, 1% ethyl-3(dimethylaminopropyl)carbodiimide (Willingham and Yamada, 1979), and dimethylsulphemidate (20 μg/ml) (Kuettel et al., 1985); and paranbenzoquinone in 0.1 M PBS (Pearse and Polak, 1975). Although antigenicity was maintained with acetone/methanol fixation, the plasticity of the frozen block was incompatible with ultracyosectioning. Therefore, no form of fixation was used before cryoprotection and the ionic strength of all solutions were maintained at or below 0.1 M to prevent the solubilization of myosin within the tissue. After cryoprotection, the tissue was rapidly frozen in liquid nitrogen.

Ultrathin (<0.1 μm) sections were cut on a cryo-system (NEW: LKB Instruments, Inc., Gaithersburg, MD) at -110°C with a tungsten-coated glass knife (Roberts, 1975). Sections were retrieved with a drop of 2.3 M sucrose containing 0.75% gelatin (type III, Sigma Chemical Co.). The gelatin reduced the surface tension of the sucrose drop as it thawed, which decreased but did not eliminate fragmentation of the unfixed tissue.

**Immunostaining**

The tissue sections were preincubated with 1% BSA in 0.1 M PBS to block nonspecific binding of the antibodies and then sequentially incubated at room temperature on a rotating table with mAb 37 at a 1:100 dilution for 45 min and goat anti-rat (GAr) IgG-5-nm gold spheres (Janssen Life Sciences Products, Piscataway, NJ) at 1:25 dilution for 45 min. All dilutions were made with 1% BSA in 0.1 M PBS. The sections were simultaneously stained and protected from drying artifact by 3% uranyl acetate in 2% methyl cellulose. The tissue was viewed with a JEOL 100CX electron microscope at 100 kV.

The level of nonspecific binding of mAb 37 was determined on tissue from PTU hypothyroid animals, which has a very low content of α-MHC. To determine if the GAr IgG gold binds directly to the tissue without the presence of the primary antibody, 2-wk baby tissue, which has a high α-MHC content, was immunostained. Although the background binding of the GAr IgG gold was very low, it was not uniformly bound to all structural components. Therefore, paired tissue sections from each experimental animal were immunostained with only the GAr IgG gold to determine the relative level of nonspecific labeling of the various regions of the sarcomere.
Figure 2. Immunolabeling of rabbit papillary muscle as a function of location within the sarcomere. Histograms of data from micrographs from 2-wk baby rabbit tissue sections immunostained with mAb 37, anti-α-MHC, and GAR IgG gold. (Top) Histogram of the number of gold grains counted for each designated region described in Fig. 1: A band, subdivided into four regions: 1, 2, 3, and M; I band; Z band; mitochondria (Mito); and intracellular regions not included in preceding categories (Other). (Center) Total area of tissue sampled for each designated region (µm²). (Bottom) Raw gold density of each sarcomere division (gr/µm²) has not been corrected for non-specific staining of GAR IgG gold.

Morphometric Analysis

A systematic sample was collected using only micrographs of longitudinally oriented sarcomeres in a non-contracted state. A minimum of nine micrographs of each tissue incubated in both mAb 37 and GAR IgG gold that contained sufficient area to yield over 1,000 grains were counted. A comparable total area was sampled from the paired section incubated only in the GAR IgG gold and yielded a low gold density.

The following classification system for the various regions of the sarcomere was used. The smallest repeating unit of the muscle, a half sarcomere, was divided into Z band, I band, and A band, which was further divided into four equally sized bins: 1, 2, 3, and M region (Fig. 1, bottom). The test lattice was placed over each micrograph to correspond with the sarcomere divisions (Fig. 1, top). At the print magnification of 48,000 used, each square of test lattice represented 0.044 µm². The total area was calculated by counting the number of squares corresponding to each region of the sarcomere and are shown as a histogram (Fig. 2).

The relative level of non-specific labeling of the GAR gold to the various regions of the sarcomere was determined by comparing the average gold densities for each region from the paired tissue sections.

Results

Thyroid Model

The effectiveness of the thyroid hormone treatment in inducing a state of hyperthyroidism in the rabbits was assessed by a decrease in body weight and by an increase of the heart weight to body weight ratio for the adult animals (Sharp et al., 1985). The average weight of the animals at the beginning of the thyroid treatment was 3.1 ± 0.1 kg. The degree of weight loss in the experimental animals was related to the length of the thyroid treatment; 1- and 2-d animals lost 70 ± 40 g (n = 3 animals; mean ± SE), while 3- and 4-d animals lost 630 ± 80 g (n = 4 animals). The heart to body weight ratios were lowest in animals injected with thyroid for 1-2 d, 1.4 ± 0.14 (n = 3); while the ratio for 3- and 4-d injected animals (1.8 ± 0.09, n = 4) was greater than the ratio found in PTU animals (1.6 ± 0.02, n = 2), but less than the value obtained for normal euthyroid animals (2.3 ± 0.03, n = 2).

Immunofluorescence

Results from the indirect immunofluorescent experiments show a homogeneous distribution of fluorescent signal throughout the population of myocytes of the left ventricle and papillary muscle. Therefore, the thyroid dose was sufficient to up-regulate expression of α-MHC in all myocytes by a similar amount and any section of tissue would thus be equally representative. The fluorescent signal is also homogeneously distributed within individual myocytes from 1-4-d thyroid-treated animals as illustrated with tissue in cross section from a 2-d animal and tissue in longitudinal section from a 4-d animal (Fig. 3, A and B, respectively).
Ultracryomicrotomy and Immunostaining

Even without the benefit of fixation, myofibrillar structures maintained their integrity and ultrastructural detail (Fig. 4 B). In the unfixed tissue the spreading artifact was not completely prevented, even though gelatin was added to the sucrose drop used to retrieve the sections (Fig. 4 A). The level of nonspecific labeling of the GAr IgG gold on the thick filaments of the PTU-treated rabbits (Fig. 5 A) is very low (7 ± 2 gr/μm², n = 8, mean ± SE). As expected, the level of nonspecific labeling is greater when both antibodies are present (22 ± 3 gr/μm², n = 8) however, this gold density is very low. This indicates there is minimal nonspecific binding of the primary antibody, mAb 37 (Fig. 5 B).

The nonspecific labeling of the secondary antibody, GAr IgG gold, was determined on tissue sections from 2-wk-old baby rabbit heart (Fig. 6). Tissue incubated in both antibodies has a very high gold density in the A band (239 ± 54 gr/μm², n = 8) while the paired tissue sections incubated only in GAr IgG gold have a very low gold density (12 ± 1 gr/μm², n = 8). Therefore, nonspecific labeling of the α-MHC antigenic site by the secondary antibody, GAr IgG gold, is minimal.

The nonspecific binding of the immunoglobulins to the sarcomere is not the same in all regions of the sarcomere. The gold densities from the paired tissue sections incubated only in GAr IgG gold showed that the values for the Z band were twice as high (18.6 ± 5.4 gr/μm², n = 14) as any other bin (7.5 ± 2.2 gr/μm² of I band, n = 14; 9.3 ± 0.5 gr/μm² of A band bin, n = 14). As the Z band is known not to contain myosin, the Z band gold density was used as an internal standard in all samples to correct for nonspecific binding of the primary antibody. An example of the level of nonspecific labeling of the GAr IgG gold obtained in these experiments is seen in Fig. 7. This figure shows the gold density obtained when the tissue was exposed to only the GAr IgG gold (solid bars) or to both ant myosin, mAb 37, and GAr IgG gold (open bars) for each of the bins within the sarcomere. Corrections were made by subtraction of half of the Z band density from the raw gold densities of the A band of that same tissue section.

The corrected gold densities for the experimental animals showed the expected increases as a function of days of thyroid treatment. Furthermore, the nascent α-MHC is incorporated throughout the length of the thick filament with the average gold density per A Band bin increasing from 38 ± 6 gr/μm² (n = 8) in the 1-d treated animals to 182 ± 60 gr/μm² (n = 8) in the 4-d treated animal. Thus the average increase of α-MHC content detected by immunogold corre-
lates with the accumulation of α-MHC found by our immunofluorescent technique and with results from pyrophosphate gels (Chizzonite et al., 1982).

There is a uniform distribution of α-MHC within the first three bins (1, 2, and 3) of the thick filament in the myocytes of the 2-wk baby as seen in Fig. 2 (bottom). As mAb 37 binds to the hinge region of the myosin molecule, a lower gold density is expected in the center (region M) where the myosin rods are oriented in an antiparallel array, thereby affording fewer antigenic sites per length of filament. A similar uniform distribution pattern is seen in tissue from the normal animals and the PTU animals (Fig. 9). Therefore, the α-MHC antigen is equally accessible to the mAb 37 throughout the length of a thick filament and any nonuniformity of gold label must be attributable to a change in α-MHC distribution.

These experiments were designed to test the pattern of incorporation of nascent α-MHC into the thick filament. A pattern of A band labeling similar to that obtained in the control tissue (see Fig. 9) would indicate that newly synthesized α-MHC could exchange equally well with existing β-MHC throughout the length of the thick filament. The electron micrographs of typical sarcomeres (Fig. 8) give the impression that a gradient exists from the I band towards the M band. The quantitative data confirms this observation. Despite the average increase of gold density with days of thyroid treatment, the end of the thick filament (bin 1) always has a greater gold density than the center of the filament (bin M). The absolute gold densities are plotted as a function of location along the length of the A band in Fig. 9 and the morphometric analysis clearly shows a nonuniform pattern. There is a greater incorporation of the nascent myosin at the end than at the center of the A band. This is most evident in the 4-d thyroid-treated animal, but is also apparent to a lesser degree at earlier times.

The data was also analyzed on a relative basis; i.e., for every animal each region within the A band was calculated as a ratio of gold density of the bin to the average A band gold density. This method will tend to diminish the systematic errors inherent in immunostaining techniques. Tissue from normal and 2-wk baby animals was used as the control and has a uniform distribution of gold density along the A band, while the thyroid-treated animals show an increased gold density above the average at the ends of the thick filament (Fig. 10). In the thyroid-treated animals the distribution was not uniform, as the gold density in bin 1 was more than 50% greater than the average value, and, correspondingly, the M

Figure 5. The level of nonspecific labeling of mAb 37, anti-α-MHC, was determined from electron micrographs of tissue from PTU-treated animals. (A) Tissue incubated with only GAR IgG gold has a low gold density. (B) Tissue incubated with both anti-α-MHC and GAR IgG gold also has a low gold density throughout the entire sarcomere. This indicates that the nonspecific labeling by the primary antibody, mAb 37 (anti-α-MHC) is minimal. Bar, 0.5 μm.
Figure 6. The level of nonspecific staining of the GAr IgG gold was determined from electron micrographs of tissue from 2-wk baby animals. (A) Tissue immunostained with only GAr IgG gold. The sparse gold labeling of this tissue indicates a low level of nonspecific labeling of the secondary antibody, GAr IgG gold. (B) Tissue immunostained with both mAb 37, anti-α-MHC, and GAr IgG gold. Note the dense gold labeling found throughout the A band and in the area of the I band. The high gold density is expected as 2-wk baby rabbit heart has a high content of α-MHC. Bar, 0.5 μm.

Figure 7. A histogram of the gold densities expressed as a function of location within the sarcomere of an animal treated with L-T3 for 4 d. Gold densities (gr/μm²) for each region of the sarcomere (see Fig. 1) were calculated from 10 micrographs. Tissue was either incubated in both anti-α-MHC, mAb 37, and GAr IgG gold (open bars) or only GAr IgG gold (solid bars). Note that though the level of nonspecific binding of the GAr IgG gold (solid bars) is low it is not uniform throughout the sarcomere as the Z band gold density is about twice as high as the other bins.

Figure 8. Immunoelectron micrographs of sarcomeres from animals injected with thyroid hormone (L-T3) for 1, 2, 3, and 4 d (A-D, respectively). Tissue has been treated with mAb 37, anti-α-MHC, and GAr IgG gold. The gold label is mainly found in the A band and increases in density with the number of days of treatment. 1, 2, 3, and M refer to subdivisions of the A band as noted at bottom of Fig. 1. I band labeling is noted in the 4-d animal, see text for discussion. Bar, 0.5 μm.
of new isomyosin into the myofibril in vivo. In the hyperthyroid model new myosin incorporates along the entire length of the thick filament with a preferential addition of the nascent α-MHC at the ends of the A band (bin 1) than in the middle (bins 3 and M).

The technique of ultracryomicrotomy was essential to this experimental model to ensure that the observed pattern of gold labeling was not an artifact due to diffusion limitations of the immunoglobulins into the tissue sections. Access to the antigenic site could be limiting in either of the alternative techniques of pre- or postembedding immunostaining. Using preembedding staining the immunoglobulin molecule must diffuse into a glycerinated myofibril structure (40 nm between thick filaments), while in postembedding staining the access to the tissue is limited by the matrix of the embedding medium. The nearly uniform distribution of gold density along the thick filament in controls, normal and 2-wk baby (Fig. 8) obtained in our experiment indicates that accessibility to antigenic sites is not a function of location along the thick filament. One limitation of the immunocytochemical technique arises from the fact that antigenicity of the tissue is lost with each of the numerous fixations tried. All incubation solutions were kept at low ionic strength in order to limit the redistribution of myosin known to occur due to solubilization of myosin at high ionic strength. Redistribution of myosin due to exchange mechanisms can not be eliminated as it is only blocked by fixation (Saad et al., 1986b).

Studying the pattern of myosin exchange in thick filaments in vivo rather than in vitro indicates the possible effect of two major components of the sarcomere, myosin-associated proteins and the thin filament, on the myosin turnover process. Though numerous myosin-associated proteins have been identified (C, H, X, M protein, and titin [Offer et al., 1973; Starr et al., 1985; Eppenberger, 1981; Maruyama et al., 1977; Wang and Ramirez-Mitchell, 1983]), surprisingly little is known about their function. It is of interest that the C zone of the A band (Sjöström et al., 1977), which contains the C protein, corresponds with those bins (3 and M) that show the lowest gold densities in the thyroid-treated animals. When the integrity of titin is altered by either stretching the sarcomere beyond overlap or by subjecting the fiber to glycercination and calcium depletion, the thick filament does not reform during thick filament reconstitution experiments (Maw and Rowe, 1986). These results suggest that myosin-associated proteins may assist in maintaining thick filament integrity and may be involved in regulation of myosin turnover.

Myosin interaction with the thin filament appears to play a critical role in both assembly and stability of the thick filament structure. Results from thick filament reconstitution experiments (Maw and Rowe, 1986) suggest that thin filaments may play a major role in the concentration of myosin in the appropriate area of the sarcomere so as to enhance thick filament assembly and alignment with the thin filament, in effect, acting as a template for the thick filament formation. The process of myosin cross-bridge interaction with actin seems to stabilize the myofibrillar structure of contracting myocytes in culture, which have more myofibrils than noncontracting myocytes (Crisona and Strohmann, 1983). Given these findings it seemed essential to maintain the cross-bridge cycling function in a model of the myosin turnover phenomena.

Most in vitro thick filament assembly models work on isolated myosin preparations exclusive of associated proteins. Although this omission limits the extent to which the findings are applicable to the in vitro condition, they nonetheless provide important facts concerning thick filament assembly and disassembly. Some myosin types (i.e., skeletal, smooth, platelet myosin) are capable of forming synthetic hybrid filaments (Wachsberger and Pepe, 1980; Pollard, 1975) while others (brain myosin and myosin from Physarum) are not (Kuczmariski and Rosenbaum, 1979; Nachmis, 1972), which implies that the structural aspects of myosin that govern self-assembly show only a small degree of myosin-type specificity. Furthermore, the different myosin types appear to be uniformly distributed throughout the hybrid filament. This is in contrast to the situation found in nematode body wall muscle. Nematode thick filaments are composed of two myosin isoforms, A and B, which are segregated to the center and ends of the thick filament, respectively (Epstein et al.,
1982). The presence of more than one myosin isoform within an individual skeletal muscle fiber (Gautier and Lowey, 1979) and within a cardiac myocyte (Samuel et al., 1983) has been detected using immunofluorescent techniques and is now an established phenomenon.

Saad et al. (1986a, c), using a fluorescence energy transfer system assay on synthetic thick filaments, have shown that myosin exchange can occur between thick filaments and is very rapid. This agrees with our result of a homogeneous distribution of the new myosin throughout the myocyte as seen at the light microscope level with indirect immunofluorescence. Over the course of the 24–96 h thyroid treatment period the new myosin is incorporated and exchanged between thick filaments, thereby presenting an appearance of uniform distribution throughout the myocytes. However, the increased accumulation at the ends of the thick filament seen with immunoelectron microscopy suggests that conditions found in vivo can prevent a homogenous distribution of the new myosin at the level of the thick filament.

The incorporation pattern of new myosin along the entire length of the thick filament is quite different from the minimal exchange of actin subunits found along the length of the thin filaments of rabbit skeletal muscle (Pardee et al., 1982). This is in contrast with the situation found in stress fibers of fibroblasts where actin incorporates throughout but has domains of slower incorporation due to the presence of actin-binding proteins (Amato and Taylor, 1986). The ability to incorporate anywhere in a filament allows for constant remodelling. Thus the pattern of myosin turnover, which allows exchange of myosin throughout the length of the thick filament, permits uninterrupted mechanical output during isoform exchange. Removal of a complete sarcomere would be detrimental to force transmission to adjacent units in series. Replacement of myosin all along the filament would be mechanically advantageous; large molecules can diffuse readily through contracting muscle (Barbosa and Da Silva, 1986).

Although with time all myosin will exchange along the thick filament, initially there is a preferential addition of the $\alpha$-MHC at the ends of the thick filament. This pattern of incorporation could be a function of the increased protein synthesis rate seen with thyroid injection. During this state of rapid growth the rates of synthesis of $\alpha$-MHC may be greater than the rates of incorporation of $\alpha$-MHC into the thick filament. This imbalance would cause an accumulation of myosin at the rate limiting step in the assembly process. One kinetic model of assembly predicts that exchange between free myosin and the thick filament would occur more frequently at the ends of the thick filament with a lower rate of exchange occurring at the center of the thick filament. This model proposes that the length regulation in thick filaments is a function of a cumulative strain mechanism and that as the filament grows longer the dissociation constant increases exponentially (Davis, 1986). Rates of disassembly of the thick filament show a similar topological pattern (Ishiwata et al., 1985).

It has been shown that actin stress fiber–like structures play a major role in directing myofibrillogenesis in cultured chick cardiac myocytes (Dlugosz et al., 1984). Work by Maw and Rowe (1985) on thick filament reconstitution in myofibrils suggests that the thin filament acts as a scaffolding template for thick filament assembly. The greatest incorporation of $\alpha$-MHC would therefore be expected in the overlap between A and I bands corresponding to bin 1 of the A Band. This is the region of the A band that shows the greatest percentage of $\alpha$-MHC incorporation in our results (Figs. 8–10).

It is also of interest to note that in growing myocytes ribosomes are located between thick filaments but excluded from the M band (Larson et al., 1973). This region corresponds to bins 1–3 of the A band in our model. The thyroid treatment is known to up-regulate both $\alpha$-MHC mRNA and rRNA (Everett et al., 1984). There is some evidence that implies translation of mRNA by the ribosome occurs in close proximity to the cellular location of that particular protein. For example, in fibroblasts, the mRNA for actin is localized near sites of actin accumulation (Lawrence and Singer, 1986).

An increased gold density in the I band was noted in two of the animal groups, the 2-wk baby and the 4-d injected animal. Any imbalance between the processes of protein synthesis and protein incorporation would be most evident in these two groups, as they are expected to have the most rapid growth rates. Though the I band is by definition that region of the sarcomere that does not contain thick filaments, it nevertheless does have as a major component titin (Wang and Ramirez-Mitchell, 1983; Maruyama et al., 1985a), which is known to bind the light meromyosin portion of myosin (Maruyama et al., 1985b). More recent work by Isaacs and Fulton (1986) has shown that nascent MHC is associated with the actin filaments of the cytoskeleton in developing chick skeletal muscle. Therefore, the nascent myosin made during the hyperthyroid period may be interacting with the proteins located in the I band. Of course the distribution of $\alpha$-MHC along the thick filament in the 2-wk baby is not expected to show the preferential accumulation of $\alpha$-MHC at the ends of the A band because the primary antibody, mAb 37, is unable to distinguish the nascent $\alpha$-MHC from the $\alpha$-MHC that is already present at high levels in the 2-wk-old baby rabbit. Furthermore, we would predict that a homogeneous distribution of $\alpha$-MHC along the A band would be present in long-term hyperthyroid animals.

The results from this experiment support the hypothesis of myosin turnover throughout the length of the thick filament in a nonsegregated manner. However, during periods of high rates of protein synthesis another aspect of the turnover process becomes evident. This involves a possible imbalance between the rates of synthesis and incorporation that results in accumulation of new myosin at the ends of the thick filament. This imbalance is a result of a factor(s) that is associated with conditions found in intact functioning myocytes.

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