To Veronika
Effects of Isoproterenol, an Adrenergic Agonist, on Resting Skeletal Muscle

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RÉSUMÉ

Cette étude démontre l’effet de l’isoprotérénol (ISO), un agoniste β-adrénergique, sur le contenu total en calcium dans le réticulum sarcoplasmique (SR), désigné [Ca\textsubscript{7}SR]. Un indicateur d’ions Ca\textsuperscript{2+} de basse affinité (K\textsubscript{D} = 2.6 mM), le tétraméthylmurexide (TMX), perméable aux membranes, a été utilisé afin d’évaluer le contenu en Ca\textsuperscript{2+} libre dans le SR, désigné [Ca\textsuperscript{2+}]\textsubscript{SR}. La fraction du TMX lié à du Ca\textsuperscript{2+} dans le SR ([Ca]\textsubscript{SR}) est proportionnel à [Ca\textsuperscript{2+}]\textsubscript{SR} laquelle peut être utilisée pour déterminer [Ca\textsubscript{7}SR].

Les résultats démontrent que 10 μM ISO augmente [Ca\textsuperscript{2+}]\textsubscript{SR} de 112 % dans les fibres musculaires maintenues à un potentiel membranaire de repos physiologique de -90 mV. Par contre, en l’absence de Ca\textsuperscript{2+} dans la solution externe, l’ISO n’augmente pas de façon significative [Ca\textsuperscript{2+}]\textsubscript{SR}, suggérant que l’ISO induit un flux de Ca\textsuperscript{2+} à travers la membrane plasmique. Cette augmentation de [Ca\textsuperscript{2+}]\textsubscript{SR} se produit avec un délai exponentiel (τ = 7.0 min) et n’est pas réversible après le retrait de l’ISO de la solution externe. Ces résultats suggèrent donc que l’ISO pourrait stimuler l’expression de certain type de canaux calcique et que ceux-ci demeurereraient actifs même après le retrait de l’agoniste β-adrénergique. Puisque l’augmentation de [Ca\textsuperscript{2+}]\textsubscript{SR} en présence d’ISO se produit au potentiel de repos physiologique, il est peu probable que des canaux calciques dépendant du voltage soient responsables de l’effet observé, même si une phosphorylation de ces canaux puisse modifier leur cinétique et donc augmenter leur probabilité d’ouverture.

De plus, nous avons observé que l’effet de l’ISO sur l’influx de Ca\textsuperscript{2+} est accru par l’hyperméilisation des fibres. Cette augmentation survient avec un délai exponentiel (τ = 1.6 min) qui ne peut être expliqué par une simple augmentation de la force motrice pour les ions...
calciques ou par le courant rectifiant dans le sens entrant. Ce résultat contredit l'hypothèse selon laquelle ISO activerait une dépendance calcique dépendante du voltage du fait que la probabilité d'ouverture de ce type de canaux augmente avec la dépolarisation membranaire.

En absence d'ISO, nous avons observé que la valeur de $f_{Ca}$ demeure la même avec ou sans $Ca^{2+}$ dans la solution externe, suggérant une absence d'influx calcique durant des conditions physiologiques de repos. Lorsque les fibres musculaires sont déplétées de 70% de leur contenu en $Ca^{2+}$, la valeur de $f_{Ca}$ n'est pas influencée par l'absence ou la présence de $Ca^{2+}$ dans le milieu extracellulaire. Ceci indique que les canaux calciques de la membrane de surface, activé par une diminution du $Ca^{2+}$ dans le RS (canaux SOC), ne sont pas impliqués directement dans la régulation du contenu en $Ca^{2+}$ dans le RS.

L'élimination de l'ISO du milieu extracellulaire produit une diminution transitoire rapide de $f_{Ca}$ qui est suivit par une ré-augmentation graduelle à une vitesse similaire de celle obtenue en présence d'ISO. Cet effet peut s'expliquer par une inversion du sens de la pompe calcique ATPase sarco-endoplasmique (SERCA), provoqué par deux facteurs : (1) l'augmentation de $[Ca^{2+}]_{SR}$ et (2) la diminution de la concentration en ATP intracellulaire ($[ATP]_i$), résultant de l'élimination de l'ISO du milieu extracellulaire.

D'autre part, nous avons observé que la stimulation adrénergique augmente le pH myoplasmique. Cette observation peut être expliquée par un effet possible de l'activation des récepteurs $\alpha$-adrénergiques, lesquels augmentent l'activité des échangeurs Na$^+$/H$^+$ via la voie des PLC et de l'IP$_3$. Ceci a pour conséquence de réduire $[H^+]$ dans le myoplasm.

Finalement, nos résultats démontrent que l'ISO augmente $[Ca_{SR}]$ au repos ainsi que l'activité des SERCA. Les résultats de cette thèse soulèvent également des questions quant au rôle physiologique des canaux SOC dans la régulation de $[Ca_{SR}]$. 

VIII
Mots clefs :

Tetraméthylmurexide, muscle squelettique, contenu en calcium, isoprotérénol, réticulum sarcoplasmique.
III. ABSTRACT

Data in this study show the effects of isoproterenol (ISO), a β-adrenergic agonist, on the total calcium content of the sarcoplasmic reticulum (SR), denoted [Ca\textsubscript{T,SR}]. Tetramethyl murexide (TMX), a membrane permeable, low affinity Ca\textsuperscript{2+} indicator was used to monitor SR Ca\textsuperscript{2+} content. The fraction of calcium-bound form of TMX in the SR (f\textsubscript{Ca}) is approximately proportional to the concentration of the free Ca\textsuperscript{2+} in the SR ([Ca\textsuperscript{2+}]\textsubscript{SR}) which in turn can be used to give [Ca\textsubscript{T,SR}].

Results show that 10 μM ISO increases [Ca\textsuperscript{2+}]\textsubscript{SR} by 112% in 30 min. in resting frog skeletal muscle. No significant increase was observed with no Ca\textsuperscript{2+} present in the external solution indicating that ISO caused a Ca\textsuperscript{2+} influx across the surface/T-system membrane. This increase in [Ca\textsuperscript{2+}]\textsubscript{SR} occurred with an exponential delay (τ = 7.0 min.) and failed to reverse after washing out ISO. These results suggest that ISO stimulates the expression of a channel-permeable to calcium during resting potentials- which would remain active or open even after washing out ISO. These results argue against the involvement of most of voltage-dependent Ca\textsuperscript{2+} channels that are gated by depolarization.

Furthermore, we showed that hyperpolarization in ISO further increases Ca\textsuperscript{2+} influx. The increase in rate of influx came on with an exponential delay (τ = 1.6 min.), and couldn’t be explained simply by the increased driving force for Ca\textsuperscript{2+} or by inward rectification. This delay also argues against the involvement of hyperpolarization-activated channels. The fact that the rate of Ca\textsuperscript{2+} flux did not decrease during hyperpolarization further supports the idea that the Ca\textsuperscript{2+} influx is not due to depolarization-activated Ca\textsuperscript{2+} channel.
In the absence of ISO, the level of $\langle C_{\text{a}} \rangle$ was the same with and without Ca$^{2+}$ in the external solution, indicative of a lack of Ca$^{2+}$ influx under resting physiological conditions. When the level of $\langle C_{\text{a}} \rangle$ was reduced to 30% of the physiological level with 20 mM EGTA, the average value of $\langle C_{\text{a}} \rangle$ was the same with or without external Ca$^{2+}$. These results thereby arguing against the involvement of store operated mechanisms in the regulation of SR Ca$^{2+}$ content in the physiological range.

The data show two well distinguished effects when ISO was removed. One is a reversible effect which showed a sudden, transient decrease in SR calcium content (termed "dip"). The "dip" can be described with a single exponential and corresponds with the rate of Ca$^{2+}$ release observed in response to a depolarization to -70 mV. The "dip" appears to require the reversal of the SR calcium pump (SERCA). After the "dip", SR calcium content rose again and reached the same rate as was observed during ISO. This steady rise in SR calcium content appeared to be the other, irreversible effect of ISO.

In addition, during the course of ISO stimulation, we observed an increase in myoplasmic pH. One possible explanation could involve the activation of $\alpha$-adrenergic receptors by ISO. The receptors activate the PLC-IP3 pathway which, in turn, enhances the Na/H exchanger and thus the removal of H$^+$ ions from the myoplasm.

In summary, the data indicate that adrenergic stimulation increases [Ca$^{2+}$]$_{SR}$ in resting fibers by activating Ca$^{2+}$ influx across the surface/T-system membrane and is consistent with the expression of an unknown Ca$^{2+}$ channel. Our results also raise doubts about whether store-operated mechanisms are involved in fibers depleted to 30% of their normal [Ca$^{2+}$]$_{SR}$. The data also suggest that SERCA is directly enhanced by ISO.
Key Words:

Tetramethyl murexide, skeletal muscle, calcium content, isoproterenol, sarcoplasmic reticulum
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V. Glossary of Abbreviations

\([\text{Ca}^{2+}]_{\text{myo}}\) myoplasmic free \(\text{Ca}^{2+}\) concentration

\([\text{Ca}^{2+}]_{\text{SR}}\) sarcoplasmic reticulum free \(\text{Ca}^{2+}\) concentration

\([\text{TMX}]_T\) total TMX concentration

"dip" exponential decrease of \(\text{fCa}\) after removing ISO

0.1EGTA, NR internal solution has 0.1 mM EGTA, external solution is NR

20EGTA, NR internal solution has 20 mM EGTA, external solution is NR

ACh acetylcholine

AChE acetylcholinesterase

AKAP A kinase anchoring protein

ATP adenosine triphosphate

CaM calmodulin

CaMK calmodulin kinase

cAMP cyclic adenosine monophosphate

CICR \(\text{Ca}^{2+}\)-induced \(\text{Ca}^{2+}\) release

Cl\(^-\)/HCO\(_3^\)\(^-\) chloride/bicarbonate exchanger

CRAC calcium release-activated calcium channel

DAG diacyl glycerol

\(D_{\text{APP}}\) apparent diffusion constant

DHPR dihydropyridine receptor

ECC excitation-contraction coupling

EGTA Ethylene glycol-bis(2-aminoethyl-ether)-N,N,N,N-tetraacetic acid

\(\text{fCa}\) fraction of TMX bound with calcium

FKBP12 FK506 binding protein

GDP, GTP guanosine diphosphate, guanosine triphosphate

GPCR G-protein coupled receptor

HCN hyperpolarization-activated cyclic nucleotide-gated channels

HPR protonated form of phenol red

HTMX protonated, neutral form of TMX
IP₃ inositol triphosphate
IP₃ROC IP₃ receptor-operated channel
ISO isoproterenol
Kₐ dissociation constant
Kₘ Michaelis constant
LVA low voltage activated channels
MCT monocarboxylate transporter
MOPS 3-[N-Morpholino]propanesulfonic acid (pH buffer)
mRNA messenger RNA
nAChR nicotinic acetylcholine receptor
NBCe Na/bicarbonate co-transporter
NR normal Ringer's solution with 1.7 mM CaCl₂
Orai Transmembrane protein 142A, store-operated regulation by STIM
pH negative logarithm of hydrogen ion concentration
PIPES Piperazine-N-N-bis[2-ethanesulfonic acid] (pH buffer)
pK negative logarithm of the dissociation constant
PKA protein kinase A
PLB phospholamban
PLC phospholipase C
PMCA plasmalemmal calcium pump
Pₒ open probability of a channel
PR phenol red
RyR ryanodine receptor
SERCA sarco/endoplasmic calcium ATP-ase
SLN sarcoplasmic reticulum
SOC store-operated channel
STIM stromal interaction molecule
TMX tetramethyl murexide
TRPC transient receptor potential channel Canonical

XVII
1. INTRODUCTION

Adrenaline was isolated and identified in 1895 by Napoleon Cybulski, a Polish physiologist. In 1896, William Bates reported the clinical effects of a substance produced by the adrenal gland in the New York Medical Journal (Bates, 1896). The active principle of suprenal extract that produces its pressor effects was isolated by the joint research of John Abel in 1899 and Jokichi Takamine in 1901. Within a few years (in 1904), it was suggested that this active principle, referred to by British physiologists as “adrenaline”, was released from sympathetic nerve terminals to act on smooth muscle cells (Bennett, 1999).

The original observations of adrenaline’s effects on the contractions of frog and mammalian muscle were made by Gruber (1914) and Orbeli (1923). Orbeli observed that - when blood adrenaline level increased due to the stimulation of the lumbar sympathetic nerve in the frog - the force of the fatigued skeletal muscle, contracting due to electrical stimulation, recovered from the weakened tension. In the ‘40s, Brown and co-workers showed that the site of action of adrenaline in fatigued nerve-muscle preparations in the rat and cat is primarily upon the muscle fiber itself (Brown et al., 1948).

In curarized frog skeletal muscle, Hutter & Loewenstein (Hutter and Loewenstein, 1955) and later Oota & Nagai (Oota and Nagai, 1977) reported a positive inotropic effect of adrenaline. This enhancement of twitch tension could be due to catecholamines acting on the actomyosin system, on the sarcoplasmic reticulum (SR) calcium pump or on the
release or influx of calcium ions. Rubio and coworkers found that the enhancement of the actomyosin complex observed in heart preparations is not present in skeletal muscle (Rubio et al., 1975).

After the exclusion of the enhancement of actomyosin complex, Gonzalez-Serratos et al. (1981) set out to examine whether adrenaline has a clear positive inotropic effect on frog skeletal muscle, and if so, what mechanisms are responsible for this effect. In split fibers, they showed that the force produced with the release of Ca$^{2+}$ from the SR by caffeine was 60-100% larger when cAMP was added to the solution; simply adding adrenaline to the interior of the cell produced no changes in tension. They speculated that adrenaline acts via cAMP, and in turn, cAMP increases the amount of Ca$^{2+}$ taken up by the SR. Furthermore they suggested that the positive inotropic effect is due to the release of this extra Ca$^{2+}$ accumulated by the stimulation of the SR Ca$^{2+}$ pump and probably by an increase in the capacity of Ca$^{2+}$ binding in the SR. Further investigating with intact fibers, they found that catecholamines produced an increase of twitch tension preferably at lower stimulation frequencies. They also showed that, after bathing the intact fiber in adrenaline for 8 minutes, the first stimulation resulted in an 80% increase of force (Fig. 1). This finding suggests that Ca$^{2+}$ entry and its accumulation in the SR occurred in the resting state.

Partially based on Gonzalez-Serratos et al. (1981) findings, Arrreola and co-workers aimed to answer whether Ca$^{2+}$ channels of frog skeletal muscle are subjected to long-term modulation by adrenaline and whether this modulation could explain twitch potentiation (Arreola et al., 1987). In intact preparation after 27 minutes of exposure to adrenaline, they measured an increased force potentiation, which was greater at high frequency
stimulation. They measured an increased Ca\(^{2+}\) current (I\(_{Ca}\)) during adrenergic stimulation, and speculated that ISO increased SR calcium content; however, they did not measure sarcoplasmic reticulum free calcium concentration ([Ca\(^{2+}\)\(_{sR}\)] directly. Since addition of nifedipine, an L-type Ca\(^{2+}\) channel inhibitor did not produce a decrease on twitch potentiation, they concluded that these Ca\(^{2+}\) channels are probably not involved. They also showed the dependence of enhanced twitch potentiation on the presence of external Ca\(^{2+}\). In addition and opposed to the finding of Gonzalez-Serratos laboratory, they did not measure an increased tension for the first stimulation of the intact fiber after 25 minutes of bathing in adrenaline (Fig. 2). They observed an increase in force potentiation only after five minutes of low frequency stimulation. They concluded that adrenaline is ineffective if no stimulation was applied.

![Image](image-url)

**Fig. 1.** Effect of L-adrenaline (1 x 10^{-6} M) on the twitch tension (upper traces) and rate of tension development and decay (lower traces). In c the fiber had not been exposed to adrenaline; in a the fiber had been in a bathing solution containing adrenaline for 8 min before it was stimulated (After Gonzalez-Serratos et al., 1981)
Fig. 2. After a control twitch, adrenaline was added to the external solution and the fiber was bathed for 25 min before the series of twitches elicited at 0.07 Hz. The last twitch is a steady-state potentiation at 5 min (After Arreola et al., 1987)

The motivation for the work of this Ph.D. thesis was based on the controversy between results reported by the above mentioned laboratories: Arreola et al. (1987) reported that twitch potentiation required a train of closely spaced action potentials and attributed this requirement to the build up of voltage activation of some type of voltage-activated Ca\(^{2+}\) channels during the train of action potentials; in contrast, Gonzalez-Serratos et al. (1981) reported that adrenaline greatly enhances the first twitch response following adrenergic stimulation which suggests that the enhanced Ca\(^{2+}\) entry occurred in the resting state. They both suggested however that the enhanced twitch potentiation is due to an increased \([\text{Ca}^{2+}]_{\text{SR}}\) resulted by adrenergic stimulation, although they never directly measured it.

1.1 Anatomy of frog skeletal muscle

The primary function of skeletal muscle is to generate force or movement in response to a physiological stimulus. Normally it transduces an electrical command into mechanical response and it has to be able to create a rapid force development and sometimes
maintain its contractile force for relatively long periods (Boron and Boulapep, 2003). To achieve this goal, muscle cells have developed highly specialized anatomical structures and functional mechanisms.

1.1.1 Neuromuscular Junction

At the neuromuscular junction or endplate region, the presynaptic axon transmits an excitatory signal to the postsynaptic muscle. The frog motor endplate in its simplest form consists of an elongated, slender nerve ending embedded in a gutter-like depression of the sarcolemma (Fig. 3, middle image). This nerve terminal contains the usual synaptic organelles, capped by a thin sheath of Schwann cell cytoplasm (Fig. 3, middle image). At the active zone, where vesicles are released from the nerve terminal, these synaptic vesicles containing acetylcholine (ACh) lie adjacent to the presynaptic membrane. The postsynaptic membrane of the muscle cell contains closely packed nicotinic ACh receptors (Fig. 3, bottom image). The extracellular matrix in the synaptic cleft contains the enzyme acetylcholinesterase (AChE) which terminates the action of ACh (Peper et al., 1974).
1.1.2 Structure of the muscle fiber

Skeletal muscle fibers are multinucleated cells formed by the fusion of elongated uninucleate cells called myoblasts. Mature fibers may be as long as the muscle of which they form part, and are normally around 50 to 100 µm in diameter in amphibians. Most of
the interior of the fiber consists of the protein filaments which constitute the contractile apparatus, grouped together in bundles called myofibrils. The myofibrils also contain the internal membrane systems of the sarcoplasmic reticulum (SR) and the T tubule system.

The myofibrils have characteristic banding patterns, and the bands on adjacent myofibrils are transversely aligned so that the whole fiber appears striated under the light microscope (Fig. 4). The two main bands are the dark, strongly birefringent A band and

![Fig. 4 Light microscope image of the frog semitendinosus muscle. The darker, more birefringent A bands and the lighter, less birefringent I bands are clearly visible. The rectangular area on the fiber represents the approximate size of transilluminated spot used for absorbance measurements. Fiber diameter is 85 μM (Photo by the author).](image)

the lighter, less birefringent I band. These bands alternate along the length of the myofibril. In the middle of the A band is a lighter region, the H zone, which is bisected by a darker line, the M line. A lighter region in the middle of the H zone, the L zone, can
sometimes be distinguished (Fig 5). Two kinds of protein filaments, called myofilaments, are major components of myofibrils. The thick filaments are called myosin and the thin filaments are called actin. The actin and myosin myofilaments form highly ordered units called sarcomeres, which are joined end to end at the Z line to form the myofibrils (Seeley et al., 2003).

![Fig. 5 The striation pattern of a vertebrate skeletal muscle fiber as seen under electron microscope (a), and its interpretation as two sets of filaments (b). Photo by H. E. Huxley. (Keynes and Aidley, 2001)](image)

Muscle contraction is described by the sliding filament theory of Huxley and Niedergerke (Huxley and Niedergerke, 1954), using interference microscopy of living muscle fibers. This theory suggests that contraction is brought about by movement of the thin filaments between the thick filaments. The sliding is caused by a series of cyclic reactions between the projections on the myosin filaments and active sites on the actin filaments. Each projection first attaches itself to the actin filament to form a cross-bridge, then pulls on it and finally releases it, moving back to attach to another site further along the actin filament. The presence of ATP and Ca\(^{2+}\) is needed for this process. The increase of myoplasmic calcium concentration ([Ca\(^{2+}\)\(_{\text{myo}}\)]) is the activator of muscle contraction. When [Ca\(^{2+}\)\(_{\text{myo}}\)] concentration increases (to about 10\(^{-6}\) M), cross bridges are formed
between the two sets of filaments; ATP is split and sliding occurs (Keynes and Aidley, 2001).

An internal membrane system, the sarcoplasmic reticulum supplies Ca\(^{2+}\) for contraction upon depolarization. The SR is a series of membrane-bound sacs between the myofibrils. Its major property is that it accumulates Ca\(^{2+}\) against a concentration gradient by means of a calcium pump which requires energy from the splitting of ATP for its activity. This pump, called the sarco-endoplasmic calcium ATP-ase (SERCA) serves to maintain the calcium ion concentration in the sarcoplasm of the living muscle at its low resting level (at about 0.1 \(\mu\)M), and to produce relaxation of the muscle after contraction.

The sarcolemma has, along its surface, much regularly arranged, tube-like invaginations called transverse, or T tubules. T tubules project into the muscle fiber and wrap around sarcomeres in the region of Z lines in frog muscle (in mammalian muscle, T tubules run in the region where actin and myosin filaments overlap). Near the T tubules, the SR is enlarged to form terminal cisternae (Peachey, 1965). A T tubule and the two adjacent terminal cisternae together are called a triad (Fig. 6). High-quality electron micrographs by Clara Franzini-Armstrong (1970) showed that, at the triads, the T tubule and its adjacent terminal cisternae are connected by an array of structures called “feet” (Fig. 7). These feet belong to the calcium-release receptor channel (RyR) anchored in the SR membrane. The T tubule membrane, on the other hand, contains particles grouped in fours (“tetrads”) that are positioned opposite the feet. These tetrads are groups of four voltage-gated calcium channels, also known as DHP receptors. All the DHP receptor tetrads are closely aligned with ryanodine receptors, but about half of the RyRs have no associated DHP receptors (Franzini-Armstrong, 1970).
Fig. 6 The internal membrane system of the frog sartorius muscle fiber. In frog, the T tubule system is located at the Z line region. (Peachey, 1965)

Fig. 7 A single triad in frog sartorius muscle. The SR and T-system membranes run parallel and are separated by an empty gap. The SR membrane is scalloped, (marked by "S") with the period of the scallops coinciding with the position of the feet. The periodically repeating feet are well visible. After Franzini-Aramstrong, (1970).
1.1.3 Types of skeletal muscle fibers

Some skeletal muscles are resistant to fatigue and are able to maintain tension for a relatively long period, although they don't have to contract rapidly. Examples are muscles that maintain body posture, such as the soleus muscle in the leg. On the other hand, some muscles are capable of contracting very rapidly. However, these “fast-twitch” muscles tend to fatigue quickly. A good example of a fast twitch muscle is the external muscles of the eye, which need to contract rapidly but don’t have to maintain contraction for a long period of time. Hence, muscle fibers are generally classified as slow twitch (type I) and fast twitch (type II), depending on their rate of force development. These fibers are also distinguished by their histologic appearance, resistance to fatigue, and biochemical composition (Seeley et al., 2003).

Slow twitch fibers are generally thinner; also appear red because of the abundance of the oxygen-binding protein myoglobin. These muscles rely on oxidative metabolism for energy, hence the high density of mitochondria. In amphibians, these muscles are called tonic muscles. The maximum shortening velocity of these muscles is lower than their fast type counterparts. The slower contraction is due to a different isoform of myosin light chain kinase (MLC), and myosin heavy chain kinase (MHC) present in these fibers, as Lutz an coworkers showed in Rana pipiens anterior tibialis muscle (Lutz et al., 1998).

Fast-twitch fibers differ among themselves with respect to fatigability and maximum velocity of contraction. In mammals, the so called type IIa fast twitch fibers are quite resistant to fatigue. Indeed, their metabolic rate is high; they contain high concentration of myoglobin and mitochondria to compensate for the increased rate of ATP hydrolysis. On the other hand, type IIb fast twitch fibers are not capable of sufficient oxidative
metabolism to sustain contraction. These fibers rely on the energy that is stored in glycogen and phosphocreatine, and are more easily fatigable. They have fewer mitochondria and lower concentrations of myoglobin and oxidative enzymes. Because of the low myoglobin content, they appear white (Boron and Boulapep, 2003). In amphibians, fast twitch fibers are classified as type 1, type 2, and type 3 fibers. This classification is based on the differences of maximum shortening velocity (type 1 being the fastest) and the various subunits of myosin molecules (Lannergren, 1987).

1.2 Excitation-contraction coupling in skeletal muscle

Skeletal muscle is excited by an all-or-nothing action potential that is propagated along the whole length of the fiber. This is followed by contraction. The process linking the two events is called the excitation-contraction coupling process.

1.2.1 End-plate potential and action potential

When an action potential arrives at the presynaptic terminal of the motor neuron it causes voltage gated Ca$^{2+}$ channels to open. Calcium ions enter the presynaptic terminal; initiate the release of the neurotransmitter, acetylcholine (ACh) from the synaptic vesicles into the synaptic cleft. Acetylcholinediffuses across the synaptic cleft and binds to nicotinic ACh-receptor channels (nAChR) on the postsynaptic muscle fiber membrane (Seeley et al., 2003). These receptors are nonselective cation channels that open when ACh binds to a specific site on the channel. The result is depolarization, known as the end-plate
potential, is produced. For the first time Fatt and Katz (Fatt and Katz, 1952) showed -in frog muscle using intracellular recording technique- how stimulation of the motor nerve affects the post synaptic membrane potential. If the end-plate potential exceeds the threshold for activating voltage gated Na\(^+\) channels, an action potential results. Generation of an action potential initiates the sequence of processes leading to contraction. (Boron and Boulapep, 2003). Once triggered, the action potential propagates along the sarcolemma and T-tubules. The depolarization of the T-tubule membrane causes the activation of L-type Ca\(^{2+}\) channels (also called dihydropyridine receptors or DHPRs). DHPRs are clustered in groups of four called tetrads. DHPRs play a pivotal role in EC-coupling since they serve as the voltage sensor for SR Ca\(^{2+}\) release (Rios and Brum, 1987). The coupling mechanism is thought to involve a physical link between the DHPR and the sarcoplasmic reticulum Ca\(^{2+}\) release channel (also called as ryanodine receptor or RyR channel).

1.2.2 Central Role of sarcoplasmic reticulum
As depicted in Fig. 6, the sarcoplasmic reticulum consists of a series of membrane-bound sacs between the myofibrils of the skeletal muscle. The volume of the SR is about 16% of that of the volume of the sarcoplasm measured in frog sartorius muscle fibers bathed in isotonic solution (Birks and Davey, 1969). In this relatively small lumen, the SR sequesters the calcium. The release of this sequestered calcium is necessary for initiating contraction upon stimulation. Hence, the SR most important property is that it accumulates Ca\(^{2+}\) against its concentration gradient (Keynes and Aidley, 2001). The protein that buffers calcium in the SR is called calsequestrin (CSQ) after the suggestion
of MacLennan. MacLennan and Wong (1971) were the first to purify this protein. Calsequestrin is a very high capacity calcium-binding protein; its molecular weight is between 44 and 65 kD (MacLennan and Wong, 1971; Meissner G et al., 1973; Campbell KP, 1980), and it has a low affinity ($K_d \approx 1 \text{ mM}$ (Pape et al., 2007)) but very high capacity for calcium. CSQ is considered to be responsible for the ability of SR to maintain a high calcium content and also maintain the necessary driving force for calcium release upon stimulation (Franzini-Armstrong et al., 1987; Pape et al., 2007; Pape and Fénelon, 2008). Pape et al. (2007) showed in cut frog muscle fibers that almost all of the calcium in the SR is bound to calsequestrin. They estimated that upon stimulation - 23 times more released calcium comes from calsequestrin as opposed to from the pool of free $\text{Ca}^{2+}$ in the SR indicative of calsequestrin importance as a buffer of $[\text{Ca}^{2+}]_{\text{SR}}$ during release.

Constantin et al., (1965) observed that calcium accumulated in the terminal sacs of sarcoplasmic reticulum in oxalate treated frog muscle. They identified these regions of the SR as the intracellular calcium sink that controls the relaxation phase of the contraction-relaxation cycle. They also suggested that this area of the SR might also be the regions from which calcium is released to trigger contraction. Indeed, calsequestrin was localized to the terminal cisternae of the SR (MacLennan and Wong, 1971; Meissner, 1975). As shown in sarcoplasmic reticulum vesicles from rabbit leg muscle, calsequestrin has a tendency to aggregate on the junctional side of the membrane (Campbell KP, 1980), suggesting a direct attachment to the junctional membrane (Franzini-Armstrong et al., 1987). Zhang et. al. and other workers later found two proteins in the junctional SR: junctin and triadin (Zhang et al., 1997; Shen et al., 2007).
They suggested that these proteins interact directly to anchor calsequestrin to ryanodine receptor and form a quaternary complex that may be required for normal operation of Ca\(^{2+}\) release.

1.2.3 Role of DHPR channels

To release Ca\(^{2+}\) upon the arrival of the action potential to the triads, the SR needs a voltage sensor which senses the depolarization of the T tubule membrane. Schneider and Chandler theorized (Schneider and Chandler, 1973) and later Brum and coworkers confirmed (Brum et al., 1988b), that there are voltage sensors present in the T tubule membrane, and their action is necessary for initiating calcium release from the SR. Indeed, one surface membrane protein was located at the junctional domains of the exterior membranes which serves as voltage sensor for the SR: the L-type calcium channel, or DHPR.

Tsien and coworkers (Tsien et al. 1988) classified L-type calcium channels according to their differences in gating, ionic conductance and pharmacology. According to these criteria, they distinguished four different L-type channels: Ca\(_{1.1}\), 1.2, 1.3, 1.4 (Fig. 8). L-type channels show high sensitivity to dihydropyridines, such as nifedipine and BAY K 8644. The responsiveness to these drugs is one of the defining criteria for separating them from the other voltage gated Ca\(^{2+}\) channels. Hence they are named dihydropyridine receptor channels (DHPR). In skeletal muscle, excitation-contraction coupling Ca\(_{1.1}\) channels play a major role as the voltage sensor. These channels are primarily voltage gated but can be modified by intracellular protein-protein interactions as well.
<table>
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<td>Deactivation rate&lt;sup&gt;f&lt;/sup&gt;</td>
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Fig. 8 Types of L-type Ca<sup>2+</sup> channels

Lower case letters in index refer to the following references from where the results were taken:

<sup>a</sup>(Tsien et al., 1988), <sup>b</sup>(Snutch et al., 1990; Birbaumer et al., 1994), <sup>c</sup>(Ertel et al., 2000),
<sup>d</sup>In 10 mM calcium
<sup>e</sup>Inactivation rate at 0 mV, 10 mM Ca or 10 mM Ba extracellular, EGTA in cell, 21 °C.
<sup>f</sup>Rate of turn-off tail current at -80 to -50 mV.
<sup>g</sup>Maximum slope conductance in 110 mM Ba.
<sup>h</sup>ω-Conotoxin GVIA from Conus geographus
<sup>i</sup>Enhancement by BAY K 8644 and inhibition by nifedipine

(after Tsien et al., 1988)

L-type calcium channels were first purified using ion-exchange chromatography by Curtis and Catterall (Curtis and Catterall, 1984) from the transverse tubule membranes of skeletal muscle. Analysis of the biochemical properties revealed that the protein has five distinct subunits α<sub>1</sub>, α<sub>2</sub>, β, δ, and γ. Takahashi et al. (1987) demonstrated in purified
dihydropyridine-sensitive calcium channels -from rabbit skeletal muscle- that these subunits are located in close proximity to each other with the principal transmembrane subunit $\alpha_1$ in association with the disulfide-linked $\alpha_2\delta$ subunit, an intracellular phosphorylated $\beta$ subunit, and a transmembrane $\gamma$ subunit (Fig. 9).

In skeletal muscle, single twitches do not require Ca$^{2+}$ from the extracellular space (Armstrong et al., 1972). It has been proposed that excitation-contraction coupling occurs via direct mechanical coupling – protein-protein interaction – between the voltage sensitive Ca$^{2+}$ channel (DHPR) and the RyR1 or $\alpha$-RyR channels in mammals and amphibians respectively (Franzini-Armstrong, 1970; Adams and Beam, 1990). During EC coupling there are voltage-driven conformational changes occurring in the DHPR that
are recorded as intramembranous charge movement that precede the channel opening (Schneider and Chandler, 1973; Garcia et al., 1994). These voltage-driven charge movements are thought to reflect of movement of the voltage sensors (DHPRs) as they change to a conformation. This change in conformation presumably gates the opening of skeletal muscle RyRs (RyR1) in the SR via a mechanical, possibly direct link between the two proteins (Brum et al., 1988a). In frog skeletal muscle, intramembranous charge movements show two well distinguishable components (Schneider and Chandler, 1973; Adrian and Peres, 1977; Hui and Chandler, 1991). The early component, Qₜₑₜ, which decays exponentially, and the late “hump”, Qₓ (see Fig. 10). Pharmacological studies suggested that Qₓ might be closely associated with Ca²⁺ release from sarcoplasmic reticulum (Huang, 1982). Using voltage clamp technique, Hui (1983) also speculated that Qₓ is linked to Ca²⁺ release from the SR. This idea that Qₓ charge is caused by movement of voltage sensors in the T tubular membrane that activate Ca²⁺ release channels in the closely apposed SR membrane, presumably via mechanical coupling was supported by the results from other labs (Hui, 1983; Vergara and Caputo, 1983; Hui and Chandler, 1991; Pape and Carrier, 2002). Nifedipine, a substance that binds to DHPRs and inhibits L-type ICa inhibits Qₓ and in addition it inhibits SR Ca²⁺ release (Huang, 1990). Using nifedipine during adrenergic stimulation however, did not change the enhanced force potentiation in frog fast twitch and rat soleus muscles (Arreola et al. 1987, Cairns and Dulhunty 1993). This result suggests that catecholamines enhance a pathway other than DHPR channels for Ca²⁺ entry.
Phosphorylation sites for cAMP-dependent protein kinase (PKA) are found on the α₁ and β subunits of DHPR in skeletal muscle cells (Curtis and Catterall, 1985; Jahn et al., 1988). Ca^{2+} flux through the channel is regulated by cAMP-dependent protein phosphorylation, (Flockerzi et al., 1986; Catterall, 2000). Ion flux studies in phospholipid vesicles showed that phosphorylation of the α₁ and β subunits can greatly increase the number of functional Ca^{2+} channels in purified preparations (Nunoki et al., 1989; Mundina-Weilenmann et al., 1991); furthermore, single-channel recordings in planar bilayer membranes show increases in open probability after phosphorylation by PKA (Flockerzi et al., 1986). As biochemical studies revealed, PKA is anchored to DHPR channel by an A kinase anchoring protein (AKAP) (Rubin CS, 1994; Gray et al., 1998). The specific anchoring molecule in skeletal muscle is AKAP15, a 15 kD protein that binds to the regulatory subunit of PKA (Fig. 9) (Gray et al., 1997). This close proximity
makes the cAMP-dependent protein kinase regulation very rapid, with observable effects within 50 ms (Johnson et al., 1994; Catterall, 2000).

In cultured skeletal muscle cells, repetitive depolarization causes a dramatic enhancement of Ca\(^{2+}\) currents (Fleig and Penner, 1996) in the critical membrane potential range near to -20 mV. This potentiation is strongly voltage dependent and also is dependent on the cAMP - PKA cascade (Sculptoreanu et al., 1993). This may result from interaction between voltage-dependent gating and phosphorylation of the Ca\(^{2+}\) channel itself (Catterall, 2000). This mechanism may greatly increase Ca\(^{2+}\) channel activity during tetanic stimulation and may play a critical role in regulation of contractile force of skeletal muscle in response to adrenaline and to the frequency of stimulation of the motor nerve (Sculptoreanu et al., 1993). Also, Arreola et al. (1987) showed an increase of L-type I\(_{\text{Ca}}\) in adrenaline in response to low frequency stimulations.

1.2.4 Role of RyR channels

When DHPRs in the T tubular membrane activate the Ca\(^{2+}\) release channels of the SR membrane (RyR channels), these channels open and Ca\(^{2+}\) will be released into the sarcoplasm. Ca\(^{2+}\) release channels in the SR membrane were named after the plant alkaloid called ryanodine. Ryanodine has a potent paralytic action on skeletal muscle. It was shown to inhibit SR Ca\(^{2+}\) release by binding with high affinity to a protein present in the SR membrane (Fleischer et al., 1985). Incorporating heavy SR proteins into planar lipid bilayer, Smith et al. (Smith et al., 1985) reported single channel recordings of Ca\(^{2+}\) release channels. Based on their results, Meissner and co-workers (Meissner et al., 1986) suggested that heavy SR vesicles indeed contain Ca\(^{2+}\) release channels.
The RyR channel consists of four polypeptide subunits of 560 kDa, normally arranged in a homo-tetrameric structure, which contains a carboxyl (C) - terminal transmembrane domain, and a large amino (N) - terminal cytoplasmic domain, the foot region (Block et al., 1988; Bhat et al., 1997).

In amphibian skeletal muscle, there are two isoforms of ryanodine receptor channels present: α-RyR and β-RyR (these isoforms are homologous with the mammalian RyR1 and RyR3). Using freeze-fracture method, α-RyR isoforms were localized in the junctional region, and β-RyR isoforms were localized at the parajunctional region of the terminal cisternae (Felder and Franzini-Armstrong, 2002). By examining micrographs from frog sartorius muscle, Franzini-Armstrong (1970) showed small projections of the SR membrane whose tips were joined to the T system membrane by some amorphous material. She called these projections and the amorphous material SR feet. It has been proposed that these feet portions of RyR channels are in mechanical contact with DHPRs (Franzini-Armstrong, 1970) and as noted above, the latter serves as voltage sensor for these coupled RyRs (Schneider and Chandler, 1973). Bianchi and Bolton (1966) first proposed then supported by Ford and Podolsky (1968) and Endo et al. (1968) that Ca\textsuperscript{2+} itself can actually trigger Ca\textsuperscript{2+} release from the SR in frog skeletal muscle. Later, Fabiato (Fabiato, 1983) observed in cardiac muscle that cardiac RyR2 isoforms can be activated by the local rise (to 1 – 10 μM) of myoplasmic calcium, a phenomenon termed Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release (CICR). Rios and Pizarro proposed that in amphibians, via their voltage sensors, the depolarization-induced Ca\textsuperscript{2+} release from α-RyRs activates nearby β-RyRs through Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release (Rios and Pizarro, 1988; Rios and Pizarro, 1991). According to this idea, calcium for CICR in skeletal muscle (in opposed to in
cardiac muscle where the source of calcium is extracellular as well) comes from the SR through the junctionally located α-RyR.

Besides the alkaloid ryanodine and calcium, RyR channels can be modulated with other substances. Ogawa and Ebashi (Ogawa and Ebashi, 1976) showed an increase of Ca$^{2+}$ release from the SR in the presence of ATP on fragmented bullfrog SR. Smith et al. (1989) using radioisotope efflux measurements also in fragmented SR vesicles observed that Mg$^{2+}$ in millimolar range blocked calcium release. In lipid bilayers, they demonstrated that calmodulin inhibits RyR channel by reducing the mean duration of channel opening, without having an effect on single-channel conductance (Smith et al., 1989).

Analysis of the RyR protein sequence in skeletal muscle revealed that it contains phosphorylation sites (Suko et al., 1993) and that protein kinase A in fact phosphorylates RyR1 channels (Reiken et al., 2003). Jayaraman et al. (1992) found that in lipid bilayers FK506 binding protein (FKBP12) co-purifies with the RYR1. By co-expressing RyR and FKBP12 in insect cells, Brillantes and coworkers (1994) demonstrated that FKBP12 modulates channel gating by increasing channels with full conductance levels, decreasing open probability ($P_o$) after caffeine activation, and increasing mean open time. Marx et al., 1998, found, that in lipid bilayers, addition of FKBP12 to RYR1 induced a process, they termed as coupled gating. They hypothesized that coupled gating provides a mechanism by which RyR1 channels that are not associated with voltage-dependent Ca$^{2+}$ channels can be regulated (Marx et al., 1998). The hypothesis suggests that the binding of four FKBP12 molecules stabilizes a conformation of the RyR that favors channel opening to the maximum conductance (Ahern et al., 1997).
Further investigating in skeletal muscle of dogs with heart failure (HF), Marks and coworkers (Reiken et al., 2003) found that long term sympathetic over-stimulation (e.g. stress that caused heart failure in their experimental animals) resulted in weakened skeletal muscle contractions. In microsomes, they identified PKA phosphorylation sites on RYR1 at Ser\(^{2843}\), and that phosphorylation causes the dissociation of FKBP12 from RyR1. Dissociation of FKBP12 increases \(P_0\) of the channel (Brillantes et al., 1994; Gaburjakova et al., 2001), increases its Ca\(^{2+}\) sensitivity (Ahern et al., 1997), and in addition, inhibits coupled gating (Marx et al., 1998). They suggested that skeletal muscle contractions in HF animals become weaker because their RyR1 channels with the released FKBP12 by PKA phosphorylation becomes “leaky” causing decrease in SR \([\text{Ca}^{2+}]\) (Reiken et al., 2003). However, this idea was not supported with the more physiological skinned (EDL) and intact (FDL) preparations of rat and mouse muscle (Blazev et al., 2001). These researchers did not measure any noticeable effect of 1-3 \(\mu\)M PKA inhibitor H-89 on action potential-mediated Ca\(^{2+}\) transients.

Taken together, there are contradicting evidences of whether or not phosphorylation can cause “leaky” SR at rest.

1.2.5 Role of SERCA1 Pump

In order to produce movements, skeletal muscle has to be able to both rapidly contract and relax. During relaxation, elevated \([\text{Ca}^{2+}]_{\text{myo}}\) has to decrease. In skeletal muscle, this occurs by the SR by the Sarco-Endoplasmic Reticulum Calcium ATP-ase (SERCA). Sarco-Endoplasmic Reticulum Calcium ATP-ase is located in the membrane of SR in skeletal muscle. This protein has a total of seven known isoforms which show great
variability in sequence and tissue expression (Andersen JP and B., 1998). SERCA1a is expressed exclusively in adult skeletal muscle, whereas the SERCA1b variant is found in the same fibers in the fetus only (Brandl et al., 1987). SERCA2a is expressed in both cardiac and slow twitch skeletal muscles and its function is similar to that of SERCA1a in fast twitch muscle. (Brandl et al., 1987), SERCA2b is a housekeeping protein involved in basal $[\text{Ca}^{2+}]_{\text{myo}}$ regulation in $\beta$-cells of the pancreas (Armstrong et al., 1972; Carafoli, 2000; Arredouani et al., 2002). The expression of SERCA3 is mostly restricted to some tissues like pancreatic $\beta$-cells (Andersen JP and B., 1998; Arredouani et al., 2002), and is weakly expressed in muscle (Carafoli, 2000).

The SERCA pump is a protein with a hydrophobic region located in the lipid bilayer, and a hydrophilic region reaching into the cytoplasm (Fig. 11) (Herbette et al., 1977). The hydrophobic region has ten transmembrane segments (M1-M10) (MacLennan et al., 1985), these segments form a cavity that has high affinity for $\text{Ca}^{2+}$ and which is responsible for the translocation of the calcium ion (MacLennan et al., 1997).

The SERCA1a is the predominant form of the pump in fast twitch skeletal muscle (Brandl et al., 1987) Its major function is to control cytosolic $\text{Ca}^{2+}$ concentration (Arredouani, 2004). Its Michaelis constant for calcium ($K_m = 0.4 \mu \text{M}$) makes this protein ideal to efficiently respond to elevated $[\text{Ca}^{2+}]_{\text{myo}}$ after $\text{Ca}^{2+}$ release from the SR (Lytton et al., 1992), and restore the elevated myoplasmic calcium concentration to resting levels, around $0.1 \mu \text{M}$.
Ca\(^{2+}\) indicates the high affinity transmembrane segments (M4, M5, M6, M8) which form a cavity for calcium translocation. ASP 351 is located on the long loop connecting the cytosolic hydrophilic and the hydrophobic region in the bilayer and involved in the hydrolysis of ATP (Arredouani, 2004).

The SERCA1 pump co-purifies with sarcolipin (SLN), a 31 amino acid protein (Odermatt et al., 1998). SLN decreases Ca\(^{2+}\) affinity of SERCA1, but increases its maximal uptake rate (Fig. 12). As Fig. 12 shows, when present at low [Ca\(^{2+}\)]\(_{\text{myo}}\), SLN inhibits Ca\(^{2+}\) uptake but at high [Ca\(^{2+}\)]\(_{\text{myo}}\) it has a stimulatory effect on uptake rates of the pump (Odermatt et al., 1998).

As opposed to phospholamban (PLB), a cardiac muscle SERCA2 regulator which can be phosphorylated during \(\beta\)-adrenergic stimulation which leads to an increase in SERCA2 activity (Slack et al., 1997), SLN does not appear to be modulated by phosphorylation (Odermatt et al., 1998). Another modulator, calmodulin kinase (CaMK) can directly phosphorylate SERCA2 (Toyofuku et al., 1994), but has no effect on SERCA1 since the latter isoform does not have the Serine 38, the phosphorylation site for CaMK (Dode et al., 1996).
Fig. 12 Effects of SLN on Ca\textsuperscript{2+} uptake activity by SERCA1
SLN decreases apparent Ca\textsuperscript{2+} affinity, thereby lowering Ca\textsuperscript{2+} uptake rates at low Ca\textsuperscript{2+} concentrations, but, at saturating Ca\textsuperscript{2+} concentrations, $V_{max}$ is stimulated. (Odermatt et al., 1998).

The lack of phosphorylation by CaMK (Dode et al., 1996), or through the cAMP-PKA pathway (Odermatt et al., 1998) on SLN-SERCA1 complex, distinguishes clearly the response of skeletal muscle from that of cardiac muscle during adrenergic stimulation. However, the phosphorylation of an unknown Ca\textsuperscript{2+} channel inhibitor or an alternate, unknown enhancing effect of adrenergic stimulation via the increased [Ca\textsuperscript{2+}]_{myo} can not be ruled out (see Discussion).

1.3 Effects of adrenergic stimulation on excitation-contraction coupling

The effects of different adrenergic agonists on EC coupling have been extensively studied (for review: Tasken and Aandahl, 2004). It has been shown that adrenergic stimulation
increases force potentiation (Gonzalez-Serratos et al., 1981; Arreola et al., 1987; Cairns SP, 1993; Sculptoreanu et al., 1993), peak tension, and sustained tension (Huerta, 1991) in frog skeletal muscle. When the muscle is incubated in ISO, it develops a greater force during contractions (Gonzalez-Serratos et al., 1981; Arreola et al., 1987). Studies with skinned skeletal muscle fibers revealed, that as opposed to cardiac cells, adrenergic stimulation has no effect on the contractile mechanism of skeletal muscle (Rubio et al., 1975; Fabiato A, 1978). Adrenaline’s positive inotropic effects on frog skeletal muscle as proposed by Gonzales et al. (1981), may be caused by 1.) increased surface Ca\(^{2+}\) influx (Schmid et al., 1985; Huerta, 1991), 2.) decreased surface Ca\(^{2+}\) efflux (Gonzalez-Serratos et al., 1981), 3.) faster recycling of Ca\(^{2+}\) within the cell (Gonzalez-Serratos et al., 1981), 4.) enhanced Ca\(^{2+}\) release from the SR (Cairns SP, 1993; Ruehr et al., 2003), or 5.) enhanced SERCA1 pump (Gonzalez-Serratos et al., 1981; Rudolf et al., 2006). Since adrenaline does not enhance the contractile mechanism in skeletal muscle (Rubio et al., 1975), these effects may eventually result in a greater Ca\(^{2+}\) release from the SR upon stimulation. The next five paragraphs briefly review the main research related to these five effects.

1.3.1 Increased Ca\(^{2+}\) influx

In cultured skeletal muscle using \(^{45}\)Ca\(^{2+}\), Schmid et al. (1985) showed that isoproterenol (ISO) increased the nitredipine-sensitive L-type \(^{45}\)Ca\(^{2+}\) influx under depolarizing conditions (K\(^+\) contractions). They also found that the effects of ISO on increased \(^{45}\)Ca\(^{2+}\) influx was additive to those of depolarization in the absence of ISO (Schmid et al., 1985). Indeed, L-type Ca\(^{2+}\) channels can be phosphorylated by PKA (Sculptoreanu et al., 1993;
Johnson et al., 1994) and phosphorylation increased Ca\(^{2+}\) transport across the sarcolemma again in response to depolarization (Johnson et al., 1994). However, the significance of these channels in the activation of skeletal muscle has been questioned, since blocking of these channels did not diminish the enhancement of contractile force of the intact EDL or soleus muscle (Arreola et al., 1987; Cairns SP, 1993; Reading SA, 2003). It is also noted that this thesis is concerned with possible ISO-induced Ca\(^{2+}\) influx in resting fibers (i.e. held at the holding potential).

1.3.2 Decreased surface Ca\(^{2+}\) efflux

Curtis (1966) demonstrated Ca\(^{2+}\) efflux in resting intact fast-twitch fibers from frog using \(^{45}\)Ca\(^{2+}\). Thereby, he established the presence of both Ca\(^{2+}\) influx and efflux in the resting steady-state condition. Gonzalez-Serratos et al. (1981) hypothesized that one of adrenaline's effects on skeletal muscle is to decrease Ca\(^{2+}\) efflux across the plasmalemma. In perfused rat heart, Patten and Clark (1983) showed that adrenergic stimulation increases \([\text{Ca}^{2+}]_{\text{myo}}\). They speculated that Ca\(^{2+}\) efflux was inhibited, and Ca\(^{2+}\) influx was probably enhanced by adrenergic mechanisms (Patten and Clark, 1983). In summary, there is no data for or against the idea that adrenergic stimulation can decrease Ca\(^{2+}\) efflux.

1.3.3 Faster Ca\(^{2+}\) recycling within the cell

Gonzalez-Serratos et al. (1981) also suggested that extra calcium release from the SR which results in faster and stronger contractions may be the result of faster recycling of Ca\(^{2+}\) within the cell. Faster recycling of Ca\(^{2+}\) is -according to their suggestion- briefer
Ca$^{2+}$ release and/or enhanced rate of pumping Ca$^{2+}$ back to the SR by the SERCA and/or faster recovery from calcium inactivation. In intact skeletal muscle fibers, Rudolf, et al. (2006) monitored the dynamics of [Ca$^{2+}$]$_{SR}$ in the tibialis anterior muscle of the living mouse during adrenergic simulation while stimulating the muscle through its motor nerve. Their results did not show any difference in the kinetics of release/reuptake in the presence or absence of isoproterenol; however they demonstrated an increased efficacy of the cycle in ISO for motor nerve stimulation (e.g. more Ca$^{2+}$ release and faster reuptake of Ca$^{2+}$ in ISO during the same time period; Rudolf et al., 2006).

1.3.4 Enhanced Ca$^{2+}$ release from the SR

Ruehr et al. (2003) showed that the caffeine-induced Ca$^{2+}$ efflux of RyR1 -expressed in CHO (Chinese hamster ovary) cells- is enhanced by β-adrenergic stimulation via cAMP-PKA phosphorylation. They speculated that PKA phosphorylation of RyR1 channels in skeletal muscle may play a role in muscle contractility. In their study from Marks’ laboratory, Wehrens et al. (2005) exposed their specimens to cardiac and respiratory stress for 21 days before examining these long term effects. “Hyperphosphorylation”, as they termed this long term effect of adrenergic stimulation, caused cardiac FKBP12.6 to detach from RyR2 and these channels become leaky. They speculated that leakiness causes the drop of resting SR [Ca$^{2+}$], with this, results in less Ca$^{2+}$ release upon stimulation, and weaker contractions, causing heart failure in their specimens (Wehrens et al., 2005; Andrew M. Bellinger, 2008). They observed that skeletal muscle contractions in these animals are more fatigable, and speculated that this was caused by hyperphosphorylation of skeletal RyR1 channels and lower [Ca$^{2+}$]$_{SR}$. On the other hand,
Rudolf et al. (2006), using yellow cameleon (D1ER), a genetically encoded ratiometric fluorescent \( \text{Ca}^{2+} \) indicator which is expressed in the SR, measured a greater drop in \([\text{Ca}^{2+}]_{\text{SR}}\) in ISO than in control experiments during single twitches and higher frequency stimulations. In contrast to the speculation above of a drop in the \([\text{Ca}^{2+}]_{\text{SR}}\), they also estimated a small increase in \([\text{Ca}^{2+}]_{\text{SR}}\) in ISO. However, their study can be criticized by the fact that their dye has a very high affinity (\(K_{\text{d(D1ER)}} = 60 \ \mu\text{M}, \) Palmer et al., 2004). This fact probably prevented them from reliably monitoring the changes in \([\text{Ca}^{2+}]_{\text{SR}}\) at physiological level (~0.6 mM at rest). Based on their results they suggested that adrenergic stimulation results in a greater \(\text{Ca}^{2+}\) release from the SR which in turn causes a greater force in muscle contraction.

1.3.5 Enhanced \(\text{Ca}^{2+}\) uptake into the SR

As described in section 1.2.5, the absence of PLB in fast twitch skeletal muscle makes phosphorylation of SERCA1a less likely (Brillantes et al., 1994; Marx et al., 1998; Gaburjakova et al., 2001; Marx et al., 2001). In studies in vivo and intact preparations Rudolf et al., (2006) experienced that during the same time period more \(\text{Ca}^{2+}\) is taken up by the SR in fibers stimulated by isoproterenol compared with controls. They suggest that cAMP/PKA may have an indirect effect on SERCA activity, mediated by PKA activation of glycogen metabolism which increases the availability of ATP (Rudolf et al., 2006). This effect may explain a higher \(\text{Ca}^{2+}\) loading of the SR upon \(\beta\)-adrenergic stimulation.

30
1.3.6 Other ways to enhance the force of twitch tension by adrenaline

It is important to note that besides increased $[Ca^{2+}]_{SR}$ there are other possibilities for ISO to enhance the force of twitch tension. Adrenergic stimulation can increase the force of twitch tension by modifying SR $Ca^{2+}$ release feedback mechanisms, without actually increasing $[Ca^{2+}]_{SR}$. I will shortly discuss the four most common sites where ISO could influence these mechanisms.

The first-discovered mechanism is calcium inactivation of $Ca^{2+}$ release, a negative feedback mechanism which rapidly inactivates the SR $Ca^{2+}$ release channel. This mechanism was first described by Baylor et al. (1983) and also showed by Schneider and Simon (1988) in voltage clamped intact frog muscle fiber. Jong et al. (1995a) further characterized this mechanism in frog cut fibers.

Jong et al. (1995b) also in frog cut skeletal muscle found a positive feedback mechanism of $Ca^{2+}$ release on the kinetics of $I_y$, the late component of intramembranous charge movement which is thought to be in close association with $Ca^{2+}$ release from the SR. They suggested that the acceleration of the ON kinetics of $I_y$ is produced by SR $Ca^{2+}$ release and the accompanying increase in free $[Ca^{2+}]$ that is expected to occur at myoplasmic sites near open RyR channels. They showed maximal effect when $[Ca^{2+}]_{SR}$ neared 300 $\mu$M, about a 1/10th of the physiological value.

Pape et al. (1996) found a strong negative feedback mechanism of $Ca^{2+}$ release on $I_y$ that first slows then essentially shuts down $I_y$. This mechanism becomes apparent when the rate of $Ca^{2+}$ release from the SR increases above a certain level.

Pape & Carrier (1998) found Ca-induced $Ca^{2+}$ release (CICR) evidenced by a decrease in release permeabition when $[Ca^{2+}]_{SR}$ decreases from 300 $\mu$M to near zero.
Results of Fenelon & Pape (2002) with depolarization to -60 mV indicated that activation of neighboring Ca\(^{2+}\) release channels are not involved so that Ca\(^{2+}\) released from an SR Ca\(^{2+}\) release channel increases Ca\(^{2+}\) release through an “autoregulatory process”. Pape, Fenelon & Carrier (2002) found a CICR mechanism consistent with recruitment of neighboring channels at more depolarized potentials (they studied -45 mV) when \([\text{Ca}^{2+}]_{\text{SR}}\) is near physiological levels. This mechanism enhances Ca\(^{2+}\) release by 7-fold or greater.

If adrenergic stimulation decreased the effect of one or both negative feedback mechanisms or enhanced one or both of the positive feedback processes the resulting greater Ca\(^{2+}\) release from the SR would cause a greater twitch tension. Since it appears that the DHPRs and RyRs might be phosphorylated by adrenergic stimulation, it seems likely that one or more of these feedback mechanisms might be affected. These effects could be additive to the increase of \([\text{Ca}^{2+}]_{\text{SR}}\) showed in this study. They also highlight the need for caution when interpreting the effects of ISO or other agents on the tension response. Namely, a change in Ca\(^{2+}\) release associated with ISO stimulation -or some other agent or condition- does not necessarily mean a change in \([\text{Ca}^{2+}]_{\text{SR}}\) as is often proposed. Also a change in \([\text{Ca}^{2+}]_{\text{SR}}\) would be not necessarily mean a corresponding change in SR Ca\(^{2+}\) release owing to the tight feed back control on Ca\(^{2+}\) release.
1.4 Effects of adrenergic stimulation on resting muscle

1.4.1 Adrenergic stimulation appears to enhance SERCA1 activity

The resting level of myoplasmic calcium in skeletal muscle is maintained by two pump proteins. These are the SERCA1 pump, and the plasmalemmal calcium pump (PMCA1) (Michalak et al., 1984; Shannon et al., 2000). These pump proteins are the most likely candidates for this job because they affinities are very close to the resting free calcium concentration in the sarcoplasm ($K_m = 0.4 \mu M$, for SERCA1 (Lytton et al., 1992), and $K_m \approx 0.2-1.0 \mu M$ for PMCA1 (Balnave and Allen, 1998)). The surface sodium-calcium exchanger (Na/CaE) is also responsible for restoring the resting levels of myoplasmic Ca$^{2+}$, but because of its low affinity for $[\text{Ca}^{2+}]_{\text{myo}}$ ($K_m = 3\mu M$), its major function is most likely the removal of Ca$^{2+}$ from the myoplasm at higher than resting concentration, e.g. during high frequency contractions (Balnave and Allen, 1998).

1.4.2 PMCA is phosphorylated by PKC

Zurini et al. (1984) showed that the Plasma-Membrane Calcium ATP-ase (PMCA) possesses an autoinhibitory domain. They demonstrated that PMCA pump can be phosphorylated by PKC in the middle of its calmodulin- (CaM) binding domain. This phosphorylation completely prevents CaM binding and lowers the affinity for Ca$^{2+}$ of the the active site ($K_a$ increased from 0.2 $\mu M$ to one around 30 $\mu M$), keeping the pump essentially inhibited (James et al., 1995). Gonzales-Serratos et al. (1981) speculated that
inhibited Ca\(^{2+}\) efflux will cause an increase in \([\text{Ca}^{2+}]_{\text{myo}}\), and this extra calcium will be removed from the myoplasm by SERCA1 and, as a result, \([\text{Ca}^{2+}]_{\text{SR}}\) will increase.

Martin et al. (1990) using light microscopic autoradiography on rat soleus and gastrocnemius muscles demonstrated the presence of α1 adrenoreceptors in slow and fast twitch skeletal muscles. Activating α1 receptors with ISO increases [PKC] through the IP3 – DAG pathway (Leeb-Lundberg et al., 1985). It is possible, they speculated that increased PKC level through adrenergic stimulation of α1 receptors (Leeb-Lundberg et al., 1985) autophosphorylates the PMCA pump causing its inhibition (James et al., 1995). This supports the possible mechanism of increasing fiber Ca\(^{2+}\) content by delaying Ca\(^{2+}\) efflux through the PMCA surface Ca\(^{2+}\) pump.

1.4.3 ISO increases myoplasmic pH

Owen (1986) showed a 0.06 pH unit alkalinization in ISO with cultured smooth muscle cells. She proposed that α1 receptor stimulation increases the IP3 level (thereby stimulating surface Na-H\(^{+}\)-exchanger (Na/HE)). Also, in frog cut skeletal muscle fibers using EGTA, Pirouzi measured a 0.05 pH unit increase of myoplasmic pH during 10-minute ISO exposure (Pirouzi, 1999).
1.5 Objectives of the study

1.5.1 Goal

Our goal was to determine if adrenergic stimulation - as previously speculated - does indeed increase \([\text{Ca}^{2+}]_{\text{SR}}\). Also, we wanted to answer the question as to whether such an increase in calcium content can occur in resting fibers (as suggested by the results of Gonzalez-Serratos et al. 1981) or whether it requires stimulation in order to activate voltage-dependent \(\text{Ca}^{2+}\) channels (as suggested by the results of Arreola et al. 1987).

1.5.2 Working hypothesis

Our working hypothesis was based on early studies with frog fast twitch skeletal muscle that showed catecholamines' positive inotropic effect on twitch tension (Oota and Nagai, 1977; Gonzalez-Serratos et al., 1981; Arreola et al., 1987). Researchers speculated, but never directly measured, that these enhancing effects are due to enhanced SR \(\text{Ca}^{2+}\) pump and thereby the increased concentration of \(\text{Ca}^{2+}\) within the SR (Gonzalez-Serratos et al., 1981), or simply the enhanced \(\text{Ca}^{2+}\) influx into the sarcoplasm from the extracellular space via a \(\text{Ca}^{2+}\) channel other than L-type (Arreola et al., 1987).

We hypothesized that adrenergic stimulation increases resting \([\text{Ca}^{2+}]_{\text{SR}}\) by increasing surface \(\text{Ca}^{2+}\) influx and in addition, it enhances SERCA. Also, we hypothesized that these enhancing effects can occur at rest.
1.5.3 Specific objectives

The objective of this thesis was to answer the following questions:

1.) Is $[\text{Ca}^{2+}]_{\text{SR}}$ increased by ISO?

It was anticipated that the answer for this objective would resolve the contradiction of possible explanations for the observed tension response of Gonzalez-Serratos et al. (1981) and Arreola et al. (1987). The former results were connected with an increase in Ca$^{2+}$ content in the resting state (enhanced tension response with 1$^{st}$ action potential) whereas the latter indicated that the enhanced tension response required multiple action potentials.

2.) Is $[\text{Ca}^{2+}]_{\text{SR}}$ (in the absence of ISO) influenced by whether or not Ca$^{2+}$ is present in the external solution?

If yes, this would indicate the presence of a significant Ca$^{2+}$ influx at rest. (N.B. that a net-zero surface Ca$^{2+}$ flux can occur with influx balanced by efflux. Eliminating influx by removing external Ca$^{2+}$ would reveal a resting efflux, if present).

3.) With partial reduction of calcium load, is $[\text{Ca}^{2+}]_{\text{SR}}$ influenced by whether or not Ca$^{2+}$ is present in the external solution?

If a store-operated mechanism –like that revealed by total depletion- is involved in regulating $[\text{Ca}^{2+}]_{\text{SR}}$ at physiological levels of $[\text{Ca}^{2+}]_{\text{SR}}$, then a Ca$^{2+}$ influx should be activated with partial depletion. This would be indicated by a greater value of $[\text{Ca}^{2+}]_{\text{SR}}$ with external Ca$^{2+}$ present.
4.) If an increase in $[Ca^{2+}]_{SR}$ by ISO is involved, is the rate of this increase influenced by hyperpolarizing the fiber (from a holding potential of -90 mV to -110 mV)?

In case if ISO increases $[Ca^{2+}]_{SR}$, are voltage-dependent $Ca^{2+}$ channels involved?

5.) Determine whether ISO effects on $[Ca^{2+}]_{SR}$ are reversible or not?

6.) Confirm whether ISO increases myoplasmic pH.
2. MATERIALS AND METHODS

2.1 Isolation and preparation of single muscle fiber obtained from the frog

In this study, we used fast twitch skeletal muscle fibers (semitendinosus) from the hind leg of the American Bullfrog (Rana catesbeiana). Bull frogs were kept at room temperature in partially water filled large terrarium containers and were fed regularly. Bull frogs were killed by decapitation and spinal cord destruction with protocols approved by the Comite d'ethique de l'experimentation animale at the Universite de Sherbrooke (protocol 126-08). Fast twitch semitendinosus and iliofibularis muscles were removed from the hind leg of the frog and then placed in isotonic, 245 mOsm Ringer's solution and kept at -2.2°C until further processing. Muscles were not used after 48 hours.

Semitendinosus or iliofibularis muscles were lightly stretched and pinned in an agar filled Petri dish under Ringer's solution. After securely pinning the muscles, the Ringer's were changed to a high-potassium, calcium-free solution (relaxing solution), which caused a transient contraction of the muscle followed by relaxation. Before isolating muscle fibers, the muscle was left for about 60 minutes until it completely relaxed.
2.2 The double Vaseline gap chamber

With the aid of a dissecting stereo microscope, a cut section of a single fiber was mechanically isolated and, after careful examination to verify visual damage, it was stretched to a sarcomere length of 3.5 – 4.0 μm and mounted on a double Vaseline-gap chamber (Fig. 13, 14) (Hille and Campbell, 1976; Pape et al., 1995; Pape et al., 2002). The cut ends of the fiber were fixed with delrin clips and the clips were held to the bottom of the end pools with Vaseline, holding the fiber at its stretched length. Vaseline seals provided electrical isolation between the central pool and the two end pools. Two Lucite covers, applied partially over the central and end pools, provided nominally complete isolation of the central and end pool solutions (Irving et al., 1987). Soon after mounting, the membrane of the fiber in the end pools were permeabilized to small molecules and ions by a 2 minute exposure to a detergent (0.01% saponin) dissolved in internal solution similar to that used for the experiment (Pape et al., 1995). The end pools were then rinsed thoroughly with the same internal solution not containing saponin. The central pool solution was changed to Ringer’s solution or 0 Ca 1 EGTA Ringer’s solution (for control experiments).

The chamber was then transferred to the experimental apparatus where electrical connections were made between the electrical apparatus and the solutions in the two end pools and in the central pool (Fig. 13, 14) with the use of agar bridges. One of the end pools, denoted by EP1, was used to measure potential (V1) and the other end pool denoted by EP2, was used to inject current (I2). The central pool (CP) was maintained at earth potential by a bath-clamp amplifier. The potential in end pool 1 was maintained at
-90 mV (V1), by passing a small holding current into end pool 2 and collecting it in the central pool. Unless indicated, V1 was always held at -90 mV. V1 was controlled by the command voltage across a 26 MΩ resistor in series with the fiber. Once on the stage of the experimental apparatus, the chamber's temperature was maintained at 14 - 16°C by its contact, with a copper block cooled by circulating water from a temperature-regulated water bath (Pape and Carrier, 1998).

![Fig. 13 Double Vaseline gap chamber.](image)

The solution that bathes the fiber is separated into three compartments by two ~300 nm long Vaseline seals (VS). EP1, the left-hand end pool used for measuring voltage V1, and EP2 the right-hand end pool used for passing current I2, as well as monitoring voltage V2. CP denotes the 500 μm long central pool, which was held at earth potential. S denotes the spot also referred to as optical recording site. The dashed lines in both end pools represent the saponin permeabilized fiber segments. The electrophysiological device is described in section 2.3. (Figure after Irving et al, 1987)
2.3 Instruments for data acquisition

During the course of an experiment a maximum of eight signals were monitored: four or six optical signals (0 – 5), and two electrical signals V1 in EP1, and I2, the current injected to the fiber in EP2 to maintain the holding potential. I2 current is the injected current that maintains a steady resting potential of the membrane (denoted I_h). I_h was used to indicate the quality of the Vaseline seals and surface/T system membrane.
2.3.1 Current clamp recording

Agar bridges, filled with the appropriate end-pool or central-pool solution, provided electrical connections between the three compartments and five small side pools (one for each electrode of V1, V0, I0, I2, V2) filled with 3 M KCl. These side pools were fitted with Ag/AgCl pellets. These pellets served as electrodes.

Before each experiment, the Ag/AgCl pellets still fitted in the side pools were bathed in 3-6% sodium hypochlorite (NaOCl – bleach) for 15 minutes. This procedure rechloridized the pellets and virtually eliminated the junction potential between the Ag/AgCl pellets and the side pool solution (3 M KCl). At the beginning of each experiment, we measured the voltage between the side pools that were connected to EP1 and CP, with a 3 M KCl agar bridge. In rare cases when the voltage was 1 mV or greater, the 3 M KCl was removed from the side pools and the rechlorization process was repeated. At the end of the experiment the measurement was repeated, so that electrode drift could be assessed. The electrode drift was generally within 1 mV.

Monitored through the left-hand end pool (EP1), fibers were held at their resting potential of -90 mV (V1, Fig. 13, 14). Currents were collected by the bath clamp in the central pool. The electrical apparatus has been described in section 2.3. Briefly, signals were filtered by a four-pole Bessel filter with a switch-selectable amplifier before and after the filter. The signals then were multiplexed and sampled by an A/D converter in an ITC16-MAC computer interface device which was connected to a MAC computer.

Since there was no action potential stimulation during these experiments, the major goal of electrical recording was to maintain the fiber at the resting potential.
2.3.2 Zeiss Axiovert 100 Inverted microscope setup

Optical measurements were performed using a Zeiss Axiovert 100 inverted microscope (Carl Zeiss Inc., Oberkochen, Germany). As Fig 15 shows, the chamber with the cut fiber was placed on the stage of the inverted microscope. When the shutter (S) was opened, the fiber was trans-illuminated by white light from a tungsten-halogen bulb (400 – 1100nm, TH). The light beam was focused on an area of the fiber (see spot S in Fig. 13, and MF) in Fig. 15) by a 20X water immersion condenser (CON) objective (ICS Acroplan Infinity-corrected objective NA 0.4; Carl Zeiss, Inc., Thornwood, NY), and the emerging light was collected by a 32X infinity-corrected (OBJ) objective (NA 0.4; Carl Zeiss Inc.), and passed through two dichroic mirrors and two beam splitters. The first dichroic mirror (505 DRLP) reflects light below 505 nm and transmits longer wavelengths. The reflected shorter wavelengths were sent through a 485(22), 470(30), or 480(30) nm band filter, focused by a lens (L1) on a photodiode (PD1). (Band filters pass light through within a range of wavelengths. For example, the 480(30) filter passes light from 465 to 495 nm). The transmitted light then passed through the second dichroic mirror (505 DRLPXR) which reflects light above 650 nm and transmits the lower wavelengths. The reflected beam passed through a 675(40) or a 690(30) band filter, and then a lens (L2), which focused the light on a photodiode (PD2). Since neither of the dyes used in our experiments absorb light at these wavelengths, this channel was used for monitoring the intrinsic change of the fiber during the experiment. The remaining light that is transmitted through the second dichroic mirror was then split by a 40/60 beam splitter and was sent through a 530(30) and a 555(50) or 560(30) band filters respectively, and picked up by two photodiodes (PD3, PD4 respectively).
Each photodiode was connected to an operational amplifier circuit, which had a 10 MΩ feedback resistor. The V2 and I2 signals passed through separate voltage followers (VF-4, World Precision Instruments, Sarasota, FL), and together with the photodiode signals, passed through separate instrumentation amplifier/filter channels. Each of the signals was filtered by a four-pole Bessel filter (824L8L-4; Frequency Devices Inc., Haverhill, MA) with its cutoff frequency set to 1 kHz. Separate switch-selectable amplifiers were placed before (gain of 1, 10, or 100) and after (gain of 1, 2, 5, or 10) the filter in each channel. The signals were multiplexed and sampled by a nominal 16-bit A/D converter in an ITC16-MAC computer interface device (Instrutech Corp., Great Neck, NY), which was connected to a Power Macintosh 7100 computer. D/A ports of the interface were also used to control the shutter and the voltage command. Stimulation and sampling protocols written in C were downloaded into the ITC-16 MAC interface, which then controlled the data acquisition during the experiment.

2.3.2.1 Calibration of the setup

This section describes how the effective extinction coefficients (ε) were determined on our setup.

The interference filters used in the experiments were calibrated for their extinction coefficients with TMX and CaTMX solutions in the above described Zeiss Axiovert 100 setup. The solutions we used for the calibration protocol were “Blank” with 150 mM KCl, 2 mM MgCl₂, 10 PIPES, pH 7.0; “0” solution with 2 mM TMX added to “Blank”; and “1” with 10 mM CaCl₂ added to the 2 mM “TMX” solution.
Fig. 15 Schematic diagram of Zeiss Axiovert 100 optical apparatus
The apparatus simultaneously measures four wavelengths (PD1 – PD4). SM, spherical mirror; TH 100W tungsten halogen lamp powered by a regulated constant-current power supply; L, collecting lens; HF, heat filter; PM, plane mirror; S, computer controlled shutter; FD, field diaphragm; CON, water immersion condenser objective; MF, fiber in the experimental chamber with glass coverslip bottom; OBJ, objective, L1 – L4 lenses focusing light onto the photodiodes PD1 – PD4; F1 – F4, bandwidth filters; 40/60 beam splitter; 505DRLP, dichroic mirror reflecting wavelengths from 449 nm - 515 nm, transmitting light with wavelengths below and above the transmitting range; 505 DRLXPR; dichroic mirror transmitting light between 530 and 670 nm, reflecting light with wavelengths below and above the transmitting range (see more details in text). Position of filters: PD1 480(30) nm, PD2 675(40) or 690(30) nm, PD3: 530(30) nm, PD4: 555(50) or 560(30) nm.
Light intensities at each wavelength \((I(\lambda))\) were recorded in "Blank", "0", and "1" solutions. From these data we calculated the absorbance \((A(\lambda))\) of the interference filters in TMX and CaTMX as follows:

\[
A_0(\lambda) = \log \frac{I_{\text{blank}}(\lambda)}{I_0(\lambda)}
\]

(2.1)

\[
A_1(\lambda) = \log \frac{I(\lambda)_{\text{blank}}}{I_1(\lambda)}
\]

(2.2)

Beer's law is given by

\[
A = \varepsilon \ast \ell \ast c
\]

(2.3)

where \(A\): absorbance, \(\varepsilon\): extinction coefficients, \(\ell\): path length of the light across the fluid, \(c\): concentration of the dye. Combining the Beer's equation for two filters, \(\lambda\) and 520 gives:

\[
\frac{A_\lambda}{A(520)} = \frac{\varepsilon_{\text{TMX}}(\lambda) \ast \ell \ast c}{\varepsilon(520) \ast \ell \ast c}
\]

(2.4)
Since the path length \((l)\) and dye concentration \((c)\) are the same, they cancel, and by rearranging:

\[
\varepsilon(\lambda)_{TMX} = \frac{A_0(\lambda) \cdot \varepsilon(520)}{A(520)}
\]  

(2.5)

which gives the extinction coefficient in TMX of the filter of \((\lambda)\) wavelength.

520 nm is the isosbestic wavelength of TMX solution. At the isosbestic wavelength the absorbance of the calcium-free and calcium-bonded forms of TMX are the same (see Fig. 16, where the two spectra intersect). The extinction coefficient at this wavelength is 15,800 M\(^{-1}\)cm\(^{-1}\) (Maylie et al., 1987).

In cases when \(Ca^{2+}\) is added to the solution, concentrations in eqn. 2.4 won’t cancel since different TMX species in different concentrations are present: free TMX ([TMX]), and TMX with \(Ca^{2+}\) bound to it ([CaTMX]). This corresponding equation is:

\[
\frac{A(\lambda)}{A(520)} = \frac{\varepsilon_{CaTMX}(\lambda) \cdot \ell \cdot [CaTMX]}{\varepsilon(520) \cdot \ell \cdot [TMX]} + \frac{\varepsilon_{TMX}(\lambda) \cdot \ell \cdot [TMX]}{\varepsilon(520) \cdot \ell \cdot [TMX]}. \tag{2.6}
\]

The extinction coefficient at wavelength \(\lambda\) in CaTMX calculated by the following steps:

\[
R_j = \frac{A_j(\lambda)}{A(520)} \tag{2.7}
\]

from eqn 2.6, and
from eqn (2.4), and

\[
R_0 = \frac{a_0(\lambda)}{A(520)} = \frac{e_{\text{TMX}}(\lambda)}{e(520)}
\]  

(2.8)

then substituting eqns. 2.7, 2.8, 2.9 into eqn. 2.6, and rearranging:

\[
R_1 = R_0 * (1 - f) + \frac{e_{\text{CaTMX}}(\lambda)}{e(520)} * f
\]  

(2.10)

and by rearranging,

\[
e_{\text{CaTMX}}(\lambda) = \frac{(R_1 - R_0(1 - f)) * e(520)}{f}
\]  

(2.11)

The concentrations of total TMX and TMX with calcium ([TMX]_r and [CaTMX], respectively) are needed for obtaining their ratio (f) in order to solve eqn. 2.11. The concentrations of these TMX species were determined by fitting (using the least squares fit method) the absorbance spectrum of CaTMX solution done on spectrophotometer.

Fig. 16 shows a TMX-CaTMX spectrum which was used for the fitting process. The isosbestic wavelength is at 520 nm, where the two spectra intersect. Absorbance values are normalized to the absorbance at the isosbestic wavelength (520 nm). The spectrum
was plotted by using absorbance data measured in every 5 nm in the range of 400 – 700 nm on the spectrophotometer.

![Graph showing absorbance spectra of CaTMX and TMX](image)

**Fig. 16 TMX absorbance spectra.**
CaTMX: TMX with calcium, TMX calcium free TMX. Absorbance values are corrected for α-Ca and normalized to the isosbestic absorbance, which is at 520 nm, where the two spectra intersect. Exp: 9 mars 2001.

For the calibration on the setup, the 520(10) filter was assumed (and verified) to be isosbestic. Its extinction coefficient was assumed to be 15,800 M⁻¹cm⁻¹, the value determined from spectrophotometer measurements for the isosbestic wavelength of 520 nm (Maylie et.al., 1987a). Absorbances with the 520(10) filter thereby served as references for the other filters (see eqn’s 2.7 and 2.8 to give $R_1$ and $R_0$, respectively).
The first column in Table 2.1 shows the bandwidth filters calibrated on the setup. The second and third columns show the calculated effective extinction coefficients, specific to the setup.

<table>
<thead>
<tr>
<th>Filters (A)</th>
<th>CTMX</th>
<th>$\varepsilon$CaTMX</th>
</tr>
</thead>
<tbody>
<tr>
<td>nm</td>
<td>M $^4$ cm$^{-1}$</td>
<td>M $^4$ cm$^{-1}$</td>
</tr>
<tr>
<td>(1)</td>
<td>7931</td>
<td>26711</td>
</tr>
<tr>
<td>480(30)</td>
<td>8071</td>
<td>27439</td>
</tr>
<tr>
<td>485(22)</td>
<td>15800</td>
<td>15800</td>
</tr>
<tr>
<td>520(10)</td>
<td>15419</td>
<td>10437</td>
</tr>
<tr>
<td>530(30)</td>
<td>11683</td>
<td>3324</td>
</tr>
<tr>
<td>555(50)</td>
<td>12495</td>
<td>2955</td>
</tr>
<tr>
<td>560(10)</td>
<td>10110</td>
<td>2406</td>
</tr>
<tr>
<td>570(30)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>675(40)</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 2.1 Extinction coefficients of the different interference filters used in the experiments
Data from filter calibration experiments: 319071, 319073. Column 1 shows the wavelengths monitored during the calibration, columns 2, 3 indicate the extinction coefficients calculated with eqns 2.1 – 2.11. $[\text{Ca}]=6.7$ mM, $[\text{TMX}]=2.21$ mM, $[\text{CaTMX}]=1.47$, $\varepsilon$Ca=0.6674, $[\text{Ca}^{2+}]=5.22$ mM,

Calculations shown in APPENDIX – C show how information about the concentrations of Ca-bound and Ca-free TMX ([CaTMX] and [TMX], respectively) is derived from the measurement-derived $A_{TMX}$ signals during an experiment using the least-squares method and the value for the extinction coefficients determined above.

2.4 Method for estimating $[\text{Ca}^{2+}]_{SR}$ with TMX

2.4.1 TMX is a reliable calcium indicator in the SR

This section explains the advantages of using TMX as calcium indicator in the SR of resting muscle. 1.) TMX has a finite but slow rate of permeability through membranes so
that it is sequestered in the SR. 2.) The $K_d$ value of TMX of 2.6 mM is significantly greater than $[Ca^{2+}]_{SR} \sim 1$ mM so that saturation is avoided thus TMX is responsive to changes in $[Ca^{2+}]_{SR}$. On the other hand, $K_d$ of TMX is not so great, that $[CaTMX]$ is reduced to an extent that it becomes difficult to measure. 3.) TMX shows little if any binding to fixed sites in the sarcoplasm (Maylie et al., 1987), also results of Pape et al. (2007) indicate a lack of binding of TMX to fixed sites in the SR. It is noted that almost all dyes do display a large degree of binding to fixed sites. 4.) Essentially no $CaTMX$ is present in the myoplasm so that all of the $CaTMX$ signal would be from the SR. The virtually zero $[CaTMX]_{myo}$ is due to the $K_d$ of TMX and the low $[Ca^{2+}]_{myo}$, the fraction of $Ca^{2+}$-bound TMX in the sarcoplasm = $[Ca^{2+}]_{myo}/([Ca^{2+}]_{myo} + K_d)$. If $[Ca^{2+}]_{myo}$ is $10^{-4}$ mM which results the $fCa_{myo}$ to be $4 \times 10^{-5}$, which is negligible. Moreover, no $Ca^{2+}$ was present in the internal solution or added to the end pools so that there should be essentially no $CaTMX$ in the sarcoplasm.

TMX, however, does have one disadvantage namely that it freely crosses SR membrane so information about the concentration of TMX in the SR ($[TMX]_{SR}$) is not readily available. One approach to deal with this problem is described in section 2.4.4.

2.4.2 Calculating $A_D(\lambda)$, the dye-related absorbance at different wavelengths

As described in Pape et. al. (2007), intrinsic absorbances were subtracted from raw absorbance ($A_{raw}$) measurements to yield the dye-related absorbances ($A_D$).

To calculate $A_D$ at a given wavelength ($\lambda$), first $A_{raw}$ was calculated:

$$A_{raw}(\lambda) = \log_{10} \frac{I_{spec, off}(\lambda)}{I(\lambda)}$$

(2.12)
where $I$ is the intensity of the light passing through a rectangular area (spot), focused on the center of the fiber, and $I_{\text{spot-off}}$ is the intensity of light when the fiber is moved out of the field.

$I_{\text{spot-off}}$ measurements were performed at the beginning and at the end of the experiment and the value of $I_{\text{spot-off}}$ for each run were determined by linear interpolation.

There was generally little change in light intensity in $I_{\text{spot-off}}$ between the beginning and the end of the experiment, so that the error due to the changing light intensity from the lamp was generally small compared to the errors discussed below.

The spot-on corrected absorbance ($A_{\text{SOC}}$) was calculated with the following formula:

$$A_{\text{SOC}}(\lambda) = A_{\text{spot-off}}(\lambda) - A_{\text{spot-on}}(\lambda)$$  \hfill (2.13)

where

$$A_{\text{spot-on}}(\lambda) = \log_{10} \frac{I_{\text{spot-off}}(\lambda)}{I_{\text{no-dye}}(\lambda)}$$  \hfill (2.14)

$A_{\text{spot-on}}(\lambda)$ is the value of the intrinsic absorbance of the fiber. $I_{\text{no-dye}}$ is the light intensity measured just before adding dye to the end pools.

Finally, the dye-related absorbance is calculated by

$$A_D(\lambda) = A_{\text{SOC}}(\lambda) - A_{\text{SOC}(\lambda_{\text{long}})}$$  \hfill (2.15)

where $A_{\text{SOC}(\lambda_{\text{long}})}$ is the spot-on corrected absorbance at long wavelength where the dye does not have detectable absorbance (675(40) nm in the majority of the experiments). (It
is important to note that the $I_{\text{spot,off}}$ values in eqns 2.12 and 2.14 are for different times, so they do not necessarily cancel when eqns 2.12 and 2.14 are substituted into eqn 2.15. The method used to obtain $A_D$ (eqn. 2.15), assumes, that any changes in intrinsic absorbance are independent of wavelength. This was verified as shown by Fig. 20 in Results.

2.4.3 Determining $fCa$ during an experiment

In this study, we monitored the absolute values of absorbances of the dye TMX at different wavelengths in order to determine the fraction of TMX complexed with Ca$^{2+}$, given by $fCa$:

$$fCa = \frac{[CaTMX]}{[TMX]} \quad (2.16)$$

The following calculation shows how $fCa$ in the following equations was determined during an experiment from the measured absorbances at wavelength $\lambda$ and at the isosbestic wavelength of 520 nm.

$$A(\lambda) = e_{TMX}(\lambda) \cdot l \cdot [TMX] + fCa(\lambda) \cdot l \cdot [CaTMX] \quad (2.17)$$

$$A(520) = e(520) \cdot l \cdot [TMX]_I \quad (2.18)$$

The ratio $R$ is calculated by:
\[ R = \frac{A(\lambda)}{A(520)} = \frac{\varepsilon_{\text{TMX}}(\lambda)}{\varepsilon(520)}(1 - f\text{Ca}) + \frac{\varepsilon_{\text{CaTMX}}}{\varepsilon(520)} f\text{Ca} \] (2.19)

and by rearranging,

\[
f\text{Ca} = \frac{\varepsilon(520) A(\lambda) - \varepsilon_{\text{TMX}}(\lambda)}{\varepsilon_{\text{CaTMX}} - \varepsilon_{\text{TMX}}(\lambda)}
\] (2.20)

2.4.4 TMX gives an estimation of total resting SR calcium \([\text{Ca}_{\text{R}}]\)

The first step towards determining \([\text{Ca}_{\text{R}}]\) was to calculate the fraction of total TMX in a fiber that is complexed with \(\text{Ca}^{2+}\), denoted by \(f\text{Ca}\) as described in the previous section (eqn's 2.17 - 2.20).

As the following steps show, \(f\text{Ca}\) gives information about \([\text{Ca}^{2+}]_{\text{SR}}\) and \([\text{Ca}_{\text{R}}]\). These steps are illustrated by the following scheme:

\[ f\text{Ca} \rightarrow f\text{Ca}_{\text{SR}} \rightarrow [\text{Ca}^{2+}]_{\text{SR}} \rightarrow [\text{Ca}_{\text{R}}] \]

The first step used the relationship between \(f\text{Ca}\) of the whole fiber given by eqn. 2.16, and the fraction of total TMX in the SR complexed with \(\text{Ca}^{2+}\) (denoted \(f\text{Ca}_{\text{SR}}\)), determined by Lamboley and Pape (2008). In their study, cut muscle fibers were equilibrated with TMX and relaxing solution internally and relaxing solution externally.

After the dye diffusion period (~ 60 minutes), optical measurements were performed to determine \(f\text{Ca}\) (Fig. 17, abscissa). 0.01% (v/v) saponin then was added to the central pool to permeabilize the sarcolemma but leave the SR membrane intact. TMX quickly diffused out of the myoplasm across the permeabilized sarcolemma into the central pool,
decreasing it's concentration to virtually zero. Optical measurements were then performed to determine \(f_{Ca_{SR}}\) (left-side ordinate in Fig. 17), plotted as filled circles. The right-side ordinate indicates the free \(Ca^{2+}\) in the SR ([\(Ca^{2+}\]_{SR}) obtained by eqn.2.21 using \(f_{Ca_{SR}}\). This gives the relationship between \(f_{Ca}\) and \(f_{Ca_{SR}}\). Once \(f_{Ca_{SR}}\) is determined, the following function was used for calculating \([Ca^{2+}]_{SR}\):

\[
[Ca^{2+}]_{SR} = \frac{f_{Ca_{SR}} * K_d}{1 - f_{Ca_{SR}}}.
\]

(2.21)

The curve in Fig 17 is the least squares best fit of eqn B6 in Appendix B to the \(f_{Ca_{SR}}\) vs. \(f_{Ca}\) data. This best fit curve can be used to estimate \([Ca^{2+}]_{SR}\) from \(f_{Ca}\) for the whole fiber.
Using the value of $[\text{Ca}^{2+}]_{\text{SR}}$ obtained with eqn. 2.21, $f_{\text{CSQ}}$ the fraction of calcium-binding sites occupied by $\text{Ca}^{2+}$ on calsequestrin from Pape et al., 2007, defined as follows:

$$f_{\text{CSQ}} = \frac{[\text{Ca}^{2+}]_{\text{SR}}^n}{[\text{Ca}^{2+}]_{\text{SR}}^n + k_{d,\text{CSQ}}}, \quad (2.22)$$

where the value of Hill coefficient $n = 2.9$, $k_{d,\text{CSQ}}$, the dissociation coefficient of CSQ for Ca$^{2+}$ is 1 mM.

The next step was to determine the total calcium content of SR ($[\text{CaT}]_{\text{SR}}$). Using the values of $[\text{Ca}^{2+}]_{\text{SR}}$ and $f_{\text{CSQ}}$ (obtained with eqn. 2.21 and 2.23, respectively) and substituting them into eqn. 2.23 below taken from Pape et al. (2007), we can calculate the total calcium content in the SR.

$$[\text{CaT}]_{\text{SR}} = [\text{Ca}^{2+}]_{\text{SR}} + f_{\text{CSQ}} \cdot Q_T + [\text{CaTMX}]_{\text{SR}}, \quad (2.23)$$

where $Q_T$ is the concentration of Ca$^{2+}$-binding sites on calsequestrin referred to SR volume. For more details and explanation of the relationship between fCa, $[\text{Ca}^{2+}]_\text{SR}$ and $[\text{CaT}]_\text{SR}$ see Fig. 23 in section 3.4 in Results.
2.5 Solutions used

2.5.1 Relaxing solution

Relaxing solution was used in Petri dish for isolating single fibers, and for mounting the cut fiber into the chamber.

Composition of Relaxing solution (in mM):

<table>
<thead>
<tr>
<th>Solution</th>
<th>K-glutamate</th>
<th>PIPES</th>
<th>EGTA</th>
<th>MgSO4·7H2O</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relaxing</td>
<td>120</td>
<td>5</td>
<td>0.1</td>
<td>1.0</td>
</tr>
</tbody>
</table>

The pH of the solution was adjusted to 7.0 with KOH, and the osmolality to 245 mOsm.

2.5.2 Internal Solutions

For the majority of the experiments we used two different internal solutions. In one we used potassium-glutamate internal solution with no calcium added (0.1 EGTA, 0Ca). In experiments where we lowered the calcium content of the fiber, the internal solutions included 20 mM EGTA (20 EGTA, 0Ca):

Composition of internal solutions (in mM):

<table>
<thead>
<tr>
<th>Solution</th>
<th>K-glutamate</th>
<th>EGTA</th>
<th>MgSO4·7H2O</th>
<th>MOPS</th>
<th>Na2ATP·3H2O</th>
<th>K2-CPx2H2O</th>
<th>K-PEP</th>
<th>Ca(OH)2</th>
</tr>
</thead>
<tbody>
<tr>
<td>K-internal</td>
<td>45</td>
<td>20</td>
<td>6.8</td>
<td>5</td>
<td>5.5</td>
<td>20</td>
<td>5</td>
<td>1.76</td>
</tr>
<tr>
<td>0 Ca 20 EGTA</td>
<td>48</td>
<td>20</td>
<td>6.8</td>
<td>5</td>
<td>5.5</td>
<td>20</td>
<td>5</td>
<td>--</td>
</tr>
<tr>
<td>0 Ca 0.1 EGTA</td>
<td>76</td>
<td>0.1</td>
<td>6.7</td>
<td>5</td>
<td>5.0</td>
<td>20</td>
<td>5</td>
<td>--</td>
</tr>
</tbody>
</table>

(K-PEP: Potassium Phospho-enol Pyruvate)
The pH was adjusted to 7.0 with KOH and the osmolality to 490 mOsm. The purpose of double osmolality was to be able to add different dyes (TMX or phenol red) to the same batch of solution. Just before the experiment, 2 mM of TMX was added and the osmolality was adjusted to 245 mOsm with water. For the experiments, where the myoplasmic pH was monitored during ISO, 1 mM phenol red was added and the osmolality was adjusted to 245 mOsm with water. For osmolality measurements Fiske osmometer (Fiske Instruments Inc.) was used.

2.5.3 External Solutions

The external solution in the center pool was Ringer's. Depending on the type of the experiment, we used either 0 Ca Ringer's (0 Ca Ringer's), or normal Ringer's (NR).

Composition of the external solutions (in mM):

<table>
<thead>
<tr>
<th>Solution</th>
<th>NaCl</th>
<th>KCl</th>
<th>Na$_2$HPO$_4$</th>
<th>NaH$_2$PO$_4$·H$_2$O</th>
<th>CaCl</th>
<th>MgCl$_2$</th>
<th>EGTA-Na</th>
</tr>
</thead>
<tbody>
<tr>
<td>NR</td>
<td>120</td>
<td>2.5</td>
<td>2.15</td>
<td>0.85</td>
<td>1.7</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>0 Ca Ringer's</td>
<td>105.5</td>
<td>2.5</td>
<td>2.15</td>
<td>0.85</td>
<td>--</td>
<td>10</td>
<td>1.0</td>
</tr>
</tbody>
</table>

The pH of the external solutions was adjusted to 7.0 with NaOH, and the osmolality to 245 mOsm.

The external solution in the center pool (1.5 ml in volume) was circulated with a speed of 3.2 ml/min using a peristaltic pump (Cole Parmer Masterflex L/S; Cole Parmer
Instrument Co., Barrington, IL, USA), exchanging the entire volume approximately 2 times per minute. The temperature of the circulating fluid was maintained at 14 - 16°C, by sending it through a hollow coil of stainless steel tubing, which was bathed in a temperature-regulated water bath directly connected to the temperature regulating system for the chamber. This approach prevents changes in fluid flux and temperature.

2.5.4 External solution change to ISO

After about 60 minutes, -the time necessary for [TMX]r to approach a leveled value at the optical recording site (see Fig. 21 in Results)- the external solution in the center pool was exchanged to one containing 10 µM isoproterenol (ISO). Five minutes before the solution change, a small amount of 100 mM ISO stock solution was made from isoproterenol-hydrochloride powder (Sigma), and then it was further diluted with external solution to the concentration of 10 µM. During the solution change the pump was switched to exchange mode. There was approximately 20 ml external solution pumped through the central pool until the pump was set back to recycling mode. The time of the exchange period was 6.5 – 7 min.

The following experiment, shown in Fig. 18, was performed to evaluate the time from the addition of the dye to the equilibration in the central pool during solution change. Due to the dead space caused by the length (160 cm) of tubing from the stopcock to the chamber (stainless steel coil and Intramedic Polyethylene Tubing, PE 205, by Becton, Dickinson and Company, Parsippany, NJ), it took 60 seconds after switching the pump to exchange mode (the beginning of the horizontal bar) for the external solution with ISO to
reach the fiber. The dye concentration reached its half maximum in approximately 80 seconds, and the near maximum, equilibrated value was measured in 2 minutes.

![Graph showing dye concentration over time](image)

**Fig. 18 Time needed for ISO to reach the fiber in the center pool**
Calibration of the peristaltic pump. Ringer's solution was used for blank; Ringer's with 2 mM TMX added was the solution denoted as DYE. The pump was switched to exchange position at the beginning of the horizontal bar. DYE reached the fiber in 60 s, reached its half maximum concentration in the central pool in 62 s, and approached its maximum concentration in 107 s. Exp: 407081.

### 2.6 Method for estimating myoplasmic pH with phenol red

The value of pH was determined by the absorbance of phenol red at the optical recording site. Fig. 22 shows the spectrum of phenol red (PR) and its protonated form (HPR). The isosbestic wavelength for phenol red is 480 nm. On the setup, this was monitored with a 480(30) interference filter.
The same dichroic mirror and filter setup of Zeiss Axiovert 100 was used as in the above mentioned experiments with TMX (see chapter 2.3.1, Fig. 15), except that the 555(50) nm filter was replaced with the 560(10) filter. The reason for this replacement is that phenol red is more sensitive to pH at this wavelength, and by using a narrower band filter we could monitor a narrow range close to the peak absorbance of unprotonated phenol red. The 675(40) nm interference filter was used to monitor the intrinsic changes of the fiber since phenol red does not absorb light at this wavelength (Fig. 19).

For determining dye related absorbances for phenol red, we used the same equations as for determining dye related TMX absorbances (eqns (2.14 – 2.15)).

![Phenol red spectrums, protonated (HPR, pH = 6), and unprotonated (PR, pH = 9.5).](image)

The isosbestic wavelength for phenol red is 480 nm, where the two waveforms intersect. This wavelength was used for monitoring phenol red concentration. The peak absorbance of PR spectrum is ~ 560 nm. Exp: Jan. 11, 2005, [HPR] = 13 μM, [PR] = 12.9 μM, in 120 mM KCl and 1 mM MOPS.

The concentration of phenol red at the optical site was estimated with Beer's law (eqn. 2.3) from the value of dye related absorbance of 480(30) (A_{D480}), and the extinction
Coefficient for phenol red at 480 was taken to be 11,000 M⁻¹cm⁻¹ (Lisman and Strong, 1979). The fractional amount of phenol red in the unprotonated form, \( f \), was estimated from

\[
f = \frac{r - r_{\text{min}}}{r_{\text{max}} - r_{\text{min}}},
\]

(2.24)

where \( r = \frac{A_{D(560)}}{A_{D(480)}} \). The values of \( r_{\text{max}} \) and \( r_{\text{min}} \) were determined with cuvettes mounted on the setup, their values were 3.21 and 0.0541, respectively (Pape and Carrier, 1998).

The value of pH was calculated then from the expression,

\[
\text{pH} = pK + \log \frac{f}{1 - f}
\]

(2.25)

where the value of pK is 7.7 (Lisman and Strong, 1979).

2.7 Statistical analyses

Student's two-tailed t test and was used to determine the statistical significance of the difference between two sets of results. The difference was considered to be significant if \( P < 0.05 \).
3. RESULTS

3.1 Intrinsic wavelength dependence

Fig. 20, panel A shows experiments carried out to evaluate whether there is wavelength dependence in the change in intrinsic absorbances during the course of experiments. In this study, the fiber was prepared, mounted, and clamped the same way as it was during a regular experiment, except there was no dye added. Upper traces (A\textsubscript{RSW}) show the raw absorbances at different wavelengths vs. time after saponin. Raw absorbances were determined by using eqn. 2.12. Lower traces (A\textsubscript{D}) show nominal dye-related absorbances from the same experiment. These were determined by eqn. 2.15. During the course of this experiment from 61 – 151 minutes, absorbance changes in the shorter wavelengths were A\textsubscript{D}(480) = -0.0080, A\textsubscript{D}(520) = -0.0115, A\textsubscript{D}(570) = -0.0062. Table 3.1 summarizes the changes in raw absorbances of the different wavelengths between 60 and 150 minutes. Column (a) gives the fiber reference; columns (1 – 4) indicate the changes in absorbance during the above given time period at the wavelengths indicated. The mean values are not significantly different from zero and not significantly different from one another. These data indicate that during the course of an experiment, there is no significant change in absorbance at the monitored wavelengths which would be caused by the change in intrinsic properties of the fiber.

Fig. 20, panel B shows an experiment which evaluates whether the addition of 10 μM ISO causes any systematic offset in intrinsic absorbances (again, as monitored by dye-related absorbance with no dye present). The conditions in this study were the same.
Fig. 20 Raw and intrinsic-corrected absorbances \( (A_\text{D}) \) in time without adding Dye, with ISO (B) and without ISO (A)

Panel A, B: Upper traces \( (A_{\text{raw}}(\lambda)) \): absorbances in four wavelengths as calculated by eqn. 2.14. Lower traces \( (A_{\text{D}}(\lambda)) \): intrinsic corrected absorbances of the same wavelengths \( (A_\text{D}) \) as calculated by eqn. 2.15. External solution Ringer's, internal solution 0 Ca K-internal.

Panel A: Exp: N14062
Panel B: the horizontal bar indicates the presence of 10 \( \mu \)M ISO in the external solution. Exp: N15062
described above, except that 10 μM ISO was added to the external solution 52 minutes after saponin treatment, indicated by the horizontal bar. Upper traces (A_{Raw}) show the raw absorbances at different wavelengths vs. time after saponin. Raw and dye related absorbances were determined as described in the previous section. Dye related absorbances -before adding ISO, and at 30 minutes in ISO- were A_D(450) = 0.0882 and 0.0896, A_D(530) = 0.0538 and 0.0565, A_D(570) = 0.0577 and 0.0593, respectively. These data do not show a significant ISO related difference in dye related absorbances. Since there is no dye present in these experiments, the non-zero A_D values in both panels are due to errors in the intrinsic correction. As will be seen later, these errors are small, -though not negligible- compared to the TMX-related absorbance values of interest.

<table>
<thead>
<tr>
<th>Exp</th>
<th>ΔA_{480}</th>
<th>ΔA_{520}</th>
<th>ΔA_{570}</th>
<th>ΔA_{690}</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>(1)</td>
<td>(2)</td>
<td>(3)</td>
<td>(4)</td>
</tr>
<tr>
<td>N14062</td>
<td>-0.0080</td>
<td>-0.0115</td>
<td>-0.0062</td>
<td>0.0240</td>
</tr>
<tr>
<td>N01062</td>
<td>0.0046</td>
<td>0.0051</td>
<td>0.0053</td>
<td>0.0011</td>
</tr>
<tr>
<td>110071</td>
<td>-0.0071</td>
<td>-0.0039</td>
<td>-0.0087</td>
<td>-0.0048</td>
</tr>
<tr>
<td>109073</td>
<td>-0.0348</td>
<td>-0.0213</td>
<td>0.0213</td>
<td></td>
</tr>
<tr>
<td>103071</td>
<td>0.0059</td>
<td>-0.0061</td>
<td>-0.0114</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>-0.0079</td>
<td>-0.0034</td>
<td>-0.0050</td>
<td>0.0060</td>
</tr>
<tr>
<td>SEM</td>
<td>0.0065</td>
<td>0.0039</td>
<td>0.0045</td>
<td>0.0063</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Exp</th>
<th>ΔA_{545}</th>
<th>ΔA_{530}</th>
<th>ΔA_{570}</th>
<th>ΔA_{690}</th>
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<tr>
<td>B</td>
<td>(1)</td>
<td>(2)</td>
<td>(3)</td>
<td>(4)</td>
</tr>
<tr>
<td>N15062</td>
<td>0.0014</td>
<td>0.0027</td>
<td>0.0016</td>
<td>0.0063</td>
</tr>
</tbody>
</table>

Table 3.1 Changes in raw absorbance by ISO (B) and without ISO (A) at different wavelengths during the course of experiments

Column (a) gives the fiber reference, columns (1 -3) give the change of dye-related absorbances at the indicated wavelengths and column 4 gives the changes in raw absorbance at 690 nm, between 60 and 160 minutes. The means are not significantly different from zero and from one another.
3.2 Diffusion of TMX in the fiber from the end pools to the optical measuring site

The most accurate measurements with TMX can be achieved when its concentration is equilibrated in the fiber's myoplasm at the optical measuring site. After equilibration, any changes in [TMX], [CaTMX] and [TMX] should be due to another controlled variable – in our case- the effect of ISO. The purpose of this experiment was to determine the time necessary for the dye to reach this steady concentration.

![Dye diffusion curve of TMX in the fiber](image)

**Fig. 21 Dye diffusion curve of TMX in the fiber**
Dye was added to the end pools 28 minutes after saponin treatment at t = 0. The curve shows a least square fit of eqn. 6 and 8 in Maylie et. al., 1987a, derived from free diffusion plus linear reversible binding (Maylie et al., 1987a; Maylie et al., 1987b). Total TMX concentration: 1.22 mM, $D_{APP} = 0.871 \times 10^{-6} \text{ cm}^2/\text{s}$, Exp: 331061.

After adding the internal solution containing TMX to the end pools, TMX began to diffuse in the fiber towards the optical recording site in the center of the central pool. Fig. 21 shows an example of the time course of dye concentration at the optical recording site.
vs. time from the addition of the dye: t = 0. The continuous curve is the least squares fit of eqn. 6 and 8 in Maylie et al., (1987a), derived from free diffusion plus linear reversible binding and/or sequestration from Maylie et al., (1987b). As indicated by the close fit of the curve to the data points, the dye diffusion within the fiber is well described by the one-dimensional diffusion equation used by Maylie et al. (1987a, eqn. 4 and 5). The path length the dye has to travel within the fiber to the optical recording site is approximately 550 μm (300 μm under the seal and 250 μm to the center of the central pool). The equilibration process between the optical site and the end pools takes about 60 minutes (Fig 21).

The multiple effects described with Fig. 22 and 23 are covered in detail in separate sections below. These sections include the increase in \(\text{fCa}^{2+}\) by ISO (section 3.3), the evaluation of the involvement of store-operated mechanisms in partially depleted fibers (section 3.4) and the effects of ISO on \([\text{Ca}^{2+}]_{\text{SR}}\) and \([\text{Car}]_{\text{SR}}\) (section 3.5).

3.3 There is an ISO-dependent \(\text{Ca}^{2+}\) influx in the steady resting fiber

To address the problem of whether calcium enters the cell in ISO at rest, two sets of conditions were prepared: one with NR and another with 0 Ca Ringer’s as external solution. In both cases, the internal solution contained 0.1 mM EGTA. Fig. 22, panel A and B show individual experiments with these conditions. The horizontal bar indicates the time period when 10 μM ISO was present in the external solution. The points clearly
show an increase in $f_{Ca}$ while in ISO when external calcium was present (Fig 22, panel A), and there is no apparent change in $f_{Ca}$ in the absence of external $Ca^{2+}$ (panel B). The thin lines in all panels in Fig. 22 are line fits to $f_{Ca}$ data points in the last 15 minutes before adding ISO and from 15 to 30 minutes in ISO. The slopes of these line fits indicate the rate of change of $f_{Ca}$ ($d/Ca/dt$). The significance of these slopes will be discussed later in this chapter. After the removal of ISO, $f_{Ca}$ quickly drops and then recovers to about the same rate of rise it had before removing ISO. Analysis and possible explanation of this "dip" in $f_{Ca}$ and subsequent recovery will be shown in section 3.8.

Table 3.2 summarizes the results from all of the experiments like that in Fig. 22. Blocks A and B show the two pairs of different experimental conditions used: with or without external calcium, respectively. In these experiments the internal solution contained 0.1 mM EGTA - resulting in normal resting $[Ca^{2+}]_{SR}$ (corresponds with Fig. 22, panel A and B). Columns 2 and 3 give the mean values of $f_{Ca}$ measured 5 minutes before adding ISO and between 28 - 30 minutes in ISO, respectively. Column 4 gives the value of the percent increase from before ISO to 28 - 30 min in ISO (% increase from column 2 to 3). The mean value of percent increase is 121% (SEM = 37%, n = 6) in fibers that contained external calcium (Block A). Panel B in Fig. 22 shows results from an experiment where no external calcium was present. There was no apparent change in ISO in this case. Similarly, the mean value of percent increase in Table 3.2 Block B, column 4 indicates no significant increase in $f_{Ca}$ with no $Ca^{2+}$ in the external solution. These results indicate that ISO causes a net influx of $Ca^{2+}$ into the fiber across the outer surface/T-system membranes.
Fig. 22 panel A, B Effects of ISO on $f_{Ca}$ in fibers with normal [Ca$^{2+}$]$_{SR}$ with NR (A) and 0 Ca Ringer's (B) external solutions. The internal solution contained 0.1 mM EGTA and 0 Ca.

The horizontal bar marks the period, when 10 μM ISO was added to the external solution. The thin lines are the least square fits to data points of the last 15 minutes before and in ISO.

A. The external solution is NR, $f_{Ca}$ before adding ISO: 0.1600, before removing ISO: 0.2780, Exp: 006061. B. The external solution is 0 Ca Ringer's, $f_{Ca}$ before adding ISO: 0.1390, before removing ISO: 0.1380, fractional increase: -0.0072, Exp: 423071.
Fig. 22 panel C, D Effects of ISO on fCa in fibers with depleted [Ca$^{2+}$]$_{\text{ER}}$ with NR (C) and 0 Ca Ringer's (D) external solutions.

The internal solution contained 20 mM EGTA and 0 Ca to decrease Ca content to the minimum. The horizontal bar marks the period when the fiber was exposed to ISO. C. the external solution is NR; during the period of ISO fCa increased more than three fold, $f_{Ca}$, last before adding and last before removing ISO: 0.0230 and 0.0830, respectively, the fractional increase is 2.6087, Exp: 226073. D. the external solution is 0 Ca Ringer's; $f_{Ca}$, last before adding and last before removing ISO: 0.0590 and 0.0793, respectively, the fractional increase is 0.3446, Exp: D11062.
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Table 3.2 Summary of experiments with normal [Ca\(^{2+}\)]\(_{\text{SR}}\) (Blocks A and B) and depleted [Ca\(^{2+}\)]\(_{\text{SR}}\) (Blocks C and D) with and without external Ca\(^{2+}\) (A, C and B, D, respectively). Column 1 shows individual experiments, columns 2 and 3 show values of fCa measured 5 minutes before adding ISO and between 28 - 30 minutes in ISO, respectively. Column 4 shows the % increase from columns 3 to 4. Columns 5 and 6 show the slope of the line fit on fCa data points 15 minute before and the last 15 minutes in ISO, respectively. (* significantly different, P < 0.05)
Columns 5 and 6 give \( \frac{dCa}{dt} \), the rate of change of \( Ca \) in the last 15 minutes before adding ISO and between 15-30 minutes in ISO, respectively. The values were obtained by fitting a line on the data points of \( Ca \) in the time periods indicated above, and shown on individual experiments in Fig. 22, panels A-D. These values will be discussed later.

For easy comparison, the mean values of \( Ca \) from Table 3.2, column 2 and 3 are illustrated in Fig. 23, in front row (5 min before ISO) and in back row (30 min in ISO), respectively. The sagittal rows A-D correspond with experiments shown in Table 3.2, rows of means A-D; yellow hue indicates fibers with normal \( [Ca^{2+}]_{SR} \) (A, B), blue hue depicts fibers with depleted (C, D) \( [Ca^{2+}]_{SR} \). In this section only rows A and B will be discussed. The main thing to note is that there is a significant increase in \( Ca \) after ~30 min in ISO when \( Ca^{2+} \) was present (row A). This is about a 2-fold increase (121%, Table 3.1, Block A, column 5). These results indicate an ISO-dependent \( Ca^{2+} \) influx in the steady resting fiber. This is one of the main experimental findings in this study.

The similar values of means of \( Ca \) in Table 3.1, Blocks A and B, column 2 also indicate that, in the absence of ISO, there is no net \( Ca^{2+} \) influx across the sarcolemma and no net \( Ca^{2+} \) leak from the SR at rest. This similarity is well represented by Fig. 23 sagittal rows A and B in the first row.
Fig. 23 Means of fCa values in fibers with close to physiological [Ca²⁺]_{SR} (sagittal rows A, B, yellow hue) and in depleted fibers (sagittal rows C, D, blue hue) The front and second rows represent data taken 5 minute before adding ISO and 28 – 30 min in ISO.

3.4 Store-operated mechanisms are not involved in regulating fiber Ca²⁺ content at about 30% of its physiological [Ca²⁺]_{SR} level

This section evaluates the involvement of store-operated mechanisms in partially depleted fibers depicted in Fig 22 and 23, panels C, D and sagittal rows C, D, respectively. Also, Table 3.2, Blocks C and D summarize data obtained from individual experiments.

To lower SR calcium content below physiological levels we added 20 mM EGTA to the calcium free internal solution. The presence of 20 mM EGTA in the internal solution decreased fCa values to about 30% of values measured in fibers with 0.1 mM EGTA.
The fact that the means of $\bar{J}_{\text{Ca}}$ values are similar in both fibers with and without external Ca\textsuperscript{2+} ($\bar{J}_{\text{Ca}} = 0.0293, 0.0257$; SEM = 0.0121, 0.0120; n = 4, 3, respectively; see Table 3.2, Blocks C, D, column 2) indicates that this level of partial depletion does not result in the involvement of store-operated mechanisms. If store-operated mechanisms were stimulated, as one might expect, $\bar{J}_{\text{Ca}}$ values should be higher in fibers with external Ca\textsuperscript{2+} present compared to those without external Ca\textsuperscript{2+}.

Data obtained by the analysis of the rate of change of $\bar{J}_{\text{Ca}}$ before the addition of ISO (Table 3.2, Blocks C and D, column 5) also indicate the absence of net Ca\textsuperscript{2+} influx in partially depleted fibers ($d\bar{J}_{\text{Ca}}/dt = 0.0001, 0.0003$; SEM = 0.0003, 0.0001; n = 4, 3 with and without external calcium, respectively – $d\bar{J}_{\text{Ca}}/dt$ values are not significantly different from zero). If Ca\textsuperscript{2+} influx were present, it would be indicated by slopes that are significantly different from zero (see Table 3.2 Blocks A, B, column 6).

3.5 Effects of ISO on $[\text{Ca}^{2+}]_{\text{SR}}$ and $[\text{Ca}_T]_{\text{SR}}$

The effects of ISO on $[\text{Ca}^{2+}]_{\text{SR}}$ and on $[\text{Ca}_T]_{\text{SR}}$ determined from the mean values of $\bar{J}_{\text{Ca}}$ data from Table 3.2, columns 2 and 3 above. The relationship between $\bar{J}_{\text{Ca}}$ and $[\text{Ca}^{2+}]_{\text{SR}}$ and detailed explanation is shown in Materials and Methods, in section 2.4.4. The relationship between $\bar{J}_{\text{Ca}}$ and $[\text{Ca}_T]_{\text{SR}}$ is determined by eqns 2.20 - 2.23 shown in Materials and Methods (Pape et al., 2007). Fig. 24 depicts this relationship using data
from experiments from this study. The abscissa shows values of $f_{Ca}$, monitored during the experiments. The ordinate shows the corresponding \([Ca_T]_{SR}\) values. The line is the best fit to the values represented by data points. The means of these values are tabulated in Table 3.3, columns 5 and 6. In this Table, column 1 gives experiments with the two different pairs of experimental conditions described in the previous section. Briefly: the first two rows A, B have 0.1 EGTA in the internal solution, the first row (A) has NR, the second (B) has 0 Ca Ringer's in the external solution. The last two rows (C, D) contain 20 mM EGTA in the internal solution, the third row (C) has NR, and the fourth (D) has 0 Ca Ringer's as external solution. Columns 2 and 3 are the means of \([Ca^{2+}]_{SR}\) values before adding ISO and the last value at 28 -30 min in ISO, respectively. These values were calculated by using the method of Pape et al. (2007), and described in the Materials and Methods section (see section 2.4.4). Columns 5 and 6 are the corresponding total

![Fig. 24 Relationship between $f_{Ca}$ and \([CaT]_{SR}\).](image)

The lowest values are obtained from fibers depleted from Ca$^{2+}$ with 20 mM EGTA in the internal solution. The highest values obtained from fibers -with 0.1 mM EGTA in the internal solution- at ~30 minutes in ISO.
calcium concentrations in the SR, calculated from the means of $[\text{Ca}^{2+}]_{\text{SR}}$ in column 3 (Last Before) and column 4 (End ISO). For calculating the $[\text{Ca}^{2+}]_{\text{SR}}$ values, eqns. B1 – B6 were used (see Appendix – B). Column 4 shows the percent increase of means of $[\text{Ca}^{2+}]_{\text{SR}}$. For calculations to obtain $[\text{CaT}]_{\text{SR}}$, we used eqn 2.23 in Materials and Methods.

One thing to note is that the % increase of [Ca2+]SR was more than two times that of [CaT]SR (253.5% vs. 112.3%). The explanation for this increase lies in the nature of the CSQ binding curve (Fig. 25).
Fig. 25 Ca-calsequestrin binding curve.
The upper dotted horizontal bar marks the range of [Ca$^{2+}$]$_{SR}$ in normal (containing 0.1 mM EGTA in the internal solution, condition A and B), the lower dotted line in depleted (with 20 mM EGTA, conditions C and D) fibers in this study (normal [Ca$^{2+}$]$_{SR}$ fibers: 0.6094 mM – 1.2936 mM, depleted: 0.1562 mM – 0.6211 mM; before ISO and at -30 min. in ISO, respectively). Modified after Fig. 5 in Pape et al. 2007.

Fig. 25 depicts the CSQ binding curve, the fraction of CSQ bound with Ca$^{2+}$ vs. [Ca$^{2+}$]$_{SR}$, modified from Pape et al., 2007. The upper dotted line indicates the range of [Ca$^{2+}$]$_{SR}$ in experiments represented by row A in Table 3.3, CSQ binding curve is the steepest in this range, indicating a greater buffering power for Ca$^{2+}$, which increases the concentration of Ca$^{2+}$-bound CSQ ([Ca-CSQ]), for smaller increase in [Ca$^{2+}$]$_{SR}$ in the SR.
The greater buffering power of CSQ in the physiological $[\text{Ca}^{2+}]_{\text{SR}}$ range seen in Fig. 25 also explains why $\Delta[\text{Ca}]_{\text{SR}}$ is greater in ISO in the physiological range of $[\text{Ca}^{2+}]_{\text{SR}}$ than in depleted fibers (15.78 mM and 6.07 mM in Table 3.3 A, column 7, and C, column 7, respectively).

### 3.6 The rate of change of $f\text{Ca}$ while in ISO

Thin lines in Fig. 22 above (described with the figure) show $df\text{Ca}/dt$ in those individual experiments. The means of these values in all experiments are shown in Table 3.2, columns 5 and 6. The fits were calculated by the best least squares fit method. A steeper slope indicates a greater $\text{Ca}^{2+}$ influx into the SR and thereby into the fiber. Fig. 26 below summarizes the individual experiments and gives the means of the slopes of line fits in both experimental conditions with and without external calcium in both normal and reduced $[\text{Ca}^{2+}]_{\text{SR}}$ fibers (see values in Table 3.2, columns 5 and 6). The solid lines represent the rate of change of $f\text{Ca}$ with $\text{Ca}^{2+}$ present and the dashed lines show values with no external calcium. The lines in ISO represent the means of slopes reached in the last 15 minutes while in ISO, they do not intend to represent the pattern of the increase of $f\text{Ca}$ (which will be considered later). Since the lines in ISO are shifted, this figure is only for illustrating the slopes.
Fig. 26 Means of change of \( fCa \) values before and in ISO in fibers with 0.1 mM (upper traces) and 20 mM (lower traces) EGTA in the internal solution. \( t_{0i} \) indicates the time of addition of ISO. The slopes were determined by the best least squares fit to the last 15 minutes of \( fCa \) values before adding ISO (\( t_{(15)} - t_{0i} \)), and the last 15 minutes in ISO (\( t_{(30)} - t_{(15)} \)). Solid line: NR external solution; dashed line: 0 Ca external solution.

Data is given in Table 3.1, columns 6 and 7.

The x axis indicates the time the fiber was exposed to ISO. The "0" on the time scale is the time of ISO addition. Negative values on the x axis indicate the time until the addition of ISO. In ISO, the rate of change of \( fCa \) increases significantly in experiments which contained \( Ca^{2+} \) in the external solution (Table 3.2, A, C, column 6). In fibers, where external \( Ca^{2+} \) was removed there was no significant change in slope while in ISO. The increased slope in ISO in fibers with NR implies an increased \( Ca^{2+} \) influx. The
interesting result with this figure is that the slope was similar in ISO with NR in both fibers with physiological and reduced $[Ca^{2+}]_{SR}$.

In summary, these results indicate that if stimulated with $\beta$-adrenergic agonists, $Ca^{2+}$ enters the cell from the extracellular space and $[Ca^{2+}]_{SR}$ increases. This $Ca^{2+}$ influx is most likely via ion channels that can open at resting potential.

3.7 The increase of $fCa$ in ISO reaches its steady rate after a delay

This section evaluates the changes of $fCa$ during the period of ISO exposure. After the addition of ISO to the NR external solution, there was a delay before the rate of increase of $fCa$ reached its final steady slope.

The data after addition of ISO in Fig. 27A is well described by an exponential and sloping line function (0.1 mM EGTA internal solution). However, the functional form of the data was not always this clear (Fig. 22A), so that the following approach was adapted to estimate the delay. The first step was to fit a line to the $fCa$ data points of the last 15 minutes before adding ISO. In Fig. 27, panel A the arrow indicates the extrapolation of this line to the time when ISO reached the fiber (Time = 0, ~65 - 70 min after dye). This point gave the value of $fCa_0$. $fCa_0$ is the predicted value of $fCa$ at the time ISO was added. The second step was to extrapolate the line fit to $fCa$ values during the last 15 minutes in ISO back to Time 0, when ISO was added (same as the time of $fCa_0$). In Fig. 27, the dotted line shows this extrapolation back to the time of addition of ISO. The
Fig. 27 A. \( fCa \) increases in the presence of 10 \( \mu \)M ISO in the external solution when Ca is present. Addition of ISO to the external solution that contains Ca\(^{2+}\) (NR), will show an exponential delay with \( \tau = 6.52 \) min until it reaches its steady slope. \( fCa_0 = 0.0660, m_0 = 0.0007, m_w = 0.0095, \) shift = 0.0079, \( \tau = 6.52 \) min; Exp: 913061 (internal solution: 0.1 EGTA).

B. Addition of Ca\(^{2+}\) to the prebathed fiber in ISO causes an instant increase in \( fCa \). The empty horizontal bar on the top indicates that 10 \( \mu \)M ISO was constantly present in the external solution. The addition of external calcium begins at the beginning of the filled horizontal line marked as “NR”, at Time = 0 min. Exp: 331081
vertical dotted-dashed line marks the difference between this latter extrapolated value and $f_{Ca0}$. This difference is referred to as the "shift".

The time constant associated with the delay ($\tau$) given by the following equation:

$$\tau = \frac{\text{shift}}{m_n - m_o}$$  \hspace{1cm} (3.1)

where $m_n$ is the slope of the fit when $t >> \tau$, $m_o$ is the slope of the line fit before adding ISO. The derivation of eqn 3.1 is given in Appendix D.

In the experiment shown in Fig. 27, panel A the shift, $f_{Ca0}$, $m_n$, and $m_o$ values were 0.0776, 0.105, 0.0012 min$^{-1}$, and 0.0050 min$^{-1}$, respectively. These values substituted into eqn 3.1 gave the value of $\tau$ of 6.5 min. Table 3.4 summarizes the results for the time constant of the delay in experiments with NR external solution with and 0.1 mM or 20 mM EGTA in the internal solution. Columns (a) in both parts of the table give the fiber reference. Columns 1 give the shift, the difference in the extrapolated value (from the fit of a sloping line) and predicted value (from line fit to data before adding ISO) of $f_{Ca}$ at the time, ISO was added. Columns 2 give the extrapolated values of $f_{Ca}$ at the time of addition of ISO ($f_{Ca0}$). Columns 3 give tau, the time constant of the delay of increase of $f_{Ca}$ in ISO, calculated by eqn. 3.1.

In summary, there is a delay with the average time constant of ~7 minutes from the time of addition of ISO until the development of a steady rise of $f_{Ca}$, and this delay is not affected by lowered $[Ca^{2+}]_{SR}$. This delay corresponds with Gonzalez-Serratos et al., (1981) observation that inotropic effect takes 8 minutes to reach its maximum level (see Discussion).
A. 0.1 EGTA internal, NR external

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B. 20 EGTA internal, NR external

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Table 3.4 Values of delay of increase of \( f_{Ca} \) in ISO with NR external solution in normal and depleted fibers

In both parts column (a) is the fiber reference, column (1) gives the difference in the extrapolated value (from exp. + slope fit) and predicted value (from line fit to data before adding ISO) of \( f_{Ca} \) at the time, ISO was added; column (2) gives \( f_{Ca} \) values at the time ISO was added, predicted from the line fit on valued before adding ISO, and finally, column (3) gives the tau of the delay in ISO.

3.7.1 Prebathing the fiber in ISO eliminates the delay in \( f_{Ca} \) increase

Another approach to investigate ISO’s effect is to expose the fiber to 0 Ca external solution with 10 μM ISO, then add Ca\(^{2+}\), and monitor the change of \( f_{Ca} \). We hypothesized, that if the delayed increase of \( f_{Ca} \) in ISO was caused by the time necessary
for the onset of enhancing mechanisms, this delay could be eliminated if ISO was already present in the external solution in advance of the addition of Ca$^{2+}$.

As we showed earlier, there is no Ca$^{2+}$ influx and increase of $fCa$ when the external solution is Ca$^{2+}$ free. Fig. 27, panel B shows a typical experiment where Ca$^{2+}$ was added to the external solution that has already contained ISO (empty bar). Ca$^{2+}$ was added twenty minutes after the addition of ISO. The minimum of 20 minutes was chosen by using the mean of tau of delay from table 3.4, part A, column 3 of ~7 min. After 20 minutes the effect of ISO is expected to reach 95% of its maximum. The upper solid horizontal bar marks the time period when Ca$^{2+}$ was present in the external solution. The results indicate a rapid rise in $fCa$. This relationship suggests that after ISO had prepared the pathways for calcium -either by phosphorylation or channel expression- without the presence of the Ca$^{2+}$ itself; the moment Ca$^{2+}$ was added, it entered the fiber across the plasma membrane and was pumped into the SR.

3.8 Removing ISO results in a sudden, transient decrease in $fCa$

This section addresses the change of values of $fCa$ after removing ISO from the external solution.

Fig. 28 shows $fCa$ values vs. time in a typical experiment when ISO was removed from the central pool. The horizontal bar marks the period of ISO presence in the external solution, and time zero on the time scale marks the time ISO was washed out. Filled and empty circles mark $fCa$ values in ISO or after ISO, respectively. When ISO was washed
out of the solution, \( fCa \) drops suddenly, then after this transient decrease, it reaches a steady positive slope similar to that measured in the last 15 minutes in ISO.

![Graph showing \( fCa \) transiently "dips" after removing ISO, then recovers its slope. The horizontal bar marks the period until ISO was present in the NR external solution. At the end of the bar, ISO was removed. \( fCa \) values decreased exponentially, then reached a steady slope, which is very similar to that in ISO (filled circles). The line marks the function least-square fitted to the data points (see text and eqn 3.3; empty circles). Slopes in ISO and after removing ISO: 0.0016 min\(^{-1}\) and 0.0023 min\(^{-1}\); \( fCa \) at the beginning of fit: 0.0989; tau of "dip": 1.47 min; Exp: O10061.

Each experiment, with NR as external solution, showed this pattern of "dip" after the removal of ISO. In cases when calcium was removed from the external solution, the phenomenon of "dip" never occurred (see Fig. 22, panel B, and D).

To evaluate this finding, we first fit a function to \( fCa \) data points after the removal of ISO. The function was an exponential plus a line given by:

\[
\Delta f_{fit} - fCa_0 = mt - C \left(1 - e^{-\frac{t}{\tau}}\right),
\]

(3.2)
where $C$ is a constant, gives the magnitude of the decrease, $m$ is the steady slope of the fit at $t = \infty$, and $t$ is time after the removal of ISO (see Appendix A for detailed information).

<table>
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<tr>
<th>Column (1)</th>
<th>$m_{\text{iso}}$</th>
<th>$m_{\text{after}}$</th>
<th>$f_{\text{Ca}(t=\text{ref})}$</th>
<th>$f_{\text{Ca Low. after}}$</th>
<th>$C$</th>
<th>$C_{f_{\text{Ca Low. after}}}$</th>
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<td>0.0500</td>
<td>0.0600</td>
<td>0.2648</td>
<td></td>
</tr>
</tbody>
</table>

$P > 0.05$

Table 3.5 Values of $f_{\text{Ca}}$ change in ISO and after removing ISO

Values obtained by an exponential and line fit to $f_{\text{Ca}}$ data in ISO and after removal of ISO (eqn. 3.3). Column (1) gives the fiber reference. For easy comparison column (2) is shown from Table 3.2, part A, are the constants obtained by the fit on $f_{\text{Ca}}$ data explained in the caption of the original table. Columns (3) and (8) are constants obtained by fitting an exponential and line to $f_{\text{Ca}}$ data after removing ISO. Column (4) is the extrapolated $f_{\text{Ca}}$ value at the time of the start of exponential decrease, column (5) is the lowest $f_{\text{Ca}}$ value after removing ISO, just before $f_{\text{Ca}}$ began to rise again to reach a constant slope. Column (6) is the constant for the exponential fit, column (7) is the ratio of column 6 and the absolute $f_{\text{Ca}}$ gain in ISO. Note, that the slopes in ISO and after the removal of ISO are not significantly different. Also note, that in column 5, the lowest value of $f_{\text{Ca}}$ is in average 12% less, than the last value in ISO, which means a 39% loss off the gained $f_{\text{Ca}}$ while in ISO (column 7).

The function was least-squares best fitted to the points after ISO. Table 3.5 gives values of the fits. Column (1) gives the fiber reference, column 2 gives the final slope of $f_{\text{Ca}}$ in ISO taken from Fig. 26, and explained there; column 3 gives the slope of the fit after removing ISO (value of $m$ in eqn. 3.2). The mean values of the two slopes are very similar, 0.0037 min$^{-1}$ and 0.0041 min$^{-1}$, respectively and not significantly different from each other. Column 4 shows the extrapolated values of $f_{\text{Ca}}$ after the removal of ISO, at the moment $f_{\text{Ca}}$ began to decrease. This value was calculated by the intersection of line before removing ISO in column 2 and the exponential fit described by eqn 3.2. Column 5 shows the lowest values of $f_{\text{Ca}}$ at the trough of the "dip". Column 6 is the magnitude of
the exponential decrease of $f\text{Ca}$ after the removal of ISO. This value gives the absolute decrease in $f\text{Ca}$. (N.B. that C is more negative than the values of $f\text{Ca}_{\text{Low After}} - f\text{Ca}_{\text{ref}}$, the minimum value, since the latter is superimposed on constant rate of increase in $f\text{Ca}$.) Column 7 gives the ratio of the absolute decrease in $f\text{Ca}$ to the value of gained $f\text{Ca}$ in ISO. From this column, the mean decrease in $f\text{Ca}$ associated with the dip was 39% of the gain of $f\text{Ca}$ associated with ISO. Column 8 shows the time constant of the exponential decrease obtained by fitting an exponential and a sloping line to $f\text{Ca}$ data after removing ISO. The mean value of 0.9133 min indicates a very rapid drop of $f\text{Ca}$ in a short time. This rapid decrease reaches its minimum value about two minutes after removing ISO then it is reversed and reaches a steady increase, $m_{\text{after}}$ (Table 3.5, column 5 and 3, respectively).

In summary, the observed "dip" and subsequent recovery indicate at least two effects of ISO, one irreversible and the other rapidly reversed. The main effect of increased Ca$^{2+}$ influx is irreversible at least in the short term of tens of minutes. This is shown by the similarity of slopes of $f\text{Ca}$ in and after ISO (0.0037 and 0.0041 min$^{-1}$, respectively) in columns 2 and 3 in table 3.5. It is important to note that the "dip" involves a significant net efflux of Ca$^{2+}$ across the SR membrane following the removal of ISO. The latter effect indicates the reversal of some effects of ISO on probably the SERCA pump (see detailed argument in Discussion).
3.9 Hyperpolarization increases Ca$^{2+}$ influx from the interstitium

To partially test whether L-type Ca$^{2+}$ channels or some other voltage dependent Ca$^{2+}$ channel is involved in the Ca$^{2+}$ influx during adrenergic stimulation, fibers were hyperpolarized for a five-minute period from the resting -90 mV to -110 mV. Fig. 29 panel A shows the plot of fCa vs. time in such an experiment with no Ca$^{2+}$ in the external solution. During the five minute hyperpolarization period, marked by the horizontal bar and the two vertical lines, fCa values did not change. Restoring the resting potential had no effect either. This result however, was expected, since it was shown in Fig. 22 panel B and D that external calcium is necessary for the increase of fCa in ISO. In the case with ISO and Ca$^{2+}$ in the external solution, the five-minute hyperpolarization resulted a large increase in fCa. This increase appeared to be exponential with an average time constant of $\tau = 1.6$ min (Table 3.6, column 4). After restoring the potential, the rate of increase gradually decreased until it reached a steady positive slope. This slope was not significantly different from that before hyperpolarization. These data suggest that increased Ca$^{2+}$ influx during adrenergic stimulation is further enhanced by hyperpolarization.
Fig. 29 Effect of hyperpolarization in ISO in absence and presence of external Ca$^{2+}$

The horizontal bars and the two vertical lines mark the 5-minute time period of hyperpolarization. In case when calcium was removed from the external solution (panel A) $f_{Ca}$ did not change during hyperpolarization. Adding calcium (panel B) caused a steady slope of increase of $f_{Ca}$ in ISO (mean: 0.0026 min$^{-1}$, SEM = 0.008, n = 4). During the 5-minute hyperpolarization $f_{Ca}$ increased with an exponential delay (mean: $t = 1.60$ min, SEM = 0.27, n = 4), reaching a slope (slope). By restoring resting potential, increase in $f_{Ca}$ decreased gradually until reaching a close to similar slope (slope after) than that before hyperpolarization (slope before). $df_{Ca}/dt$ is the time derivative of $f_{Ca}$ traces. Panel A: Exp: 310081, panel B: Exp: 303083.
Table 3.6 Effect of hyperpolarization on $f_{Ca}$ increase in ISO

The mean steady slope of $f_{Ca}$ increase during hyperpolarization is significantly greater than before or after at -90 mV (columns 3 and 2, 5, respectively). Tau (4) was obtained by $f_{Ca}$ values determined by fitting a line and extrapolated back to the time of the beginning of hyperpolarization, and the actual $f_{Ca}$ values measured at that time, and using eqns. 3.7 – 3.9.

* P < 0.05

The solid line in Fig. 29 panel B shows the time derivative of $f_{Ca}$ values. An important thing to note is that there was no decrease in $df_{Ca}/dt$ at the start of the hyperpolarization which would have been indicative of the involvement of a voltage-dependent Ca$^{2+}$ channel. On the other hand, when the holding potential was restored to -90 mV, $df_{Ca}/dt$ changed direction, began to decrease with a close to similar rate as that of its increase during hyperpolarization. Thin lines are the line fits to the data points 15 minutes before (slope before), the last three data points during hyperpolarization (slope) and between 10 – 20 minutes after restoring the holding potential to -90 mV (slope after).

Table 3.6 shows the means of slopes of lines fitted on $f_{Ca}$ values (see the slopes also in Fig. 29) in a 15-minute period before, last 3 data points during, and between 10 – 20 min. after hyperpolarization, when it reached the steady slope (columns 2,3 and 5, respectively), also the average of time constants (column 4). The time constant was calculated by using eqn 3.1.

<table>
<thead>
<tr>
<th></th>
<th>-90 mV</th>
<th>-110 mV</th>
<th>-90 mV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Slope</td>
<td>Slope</td>
<td>Slope</td>
</tr>
<tr>
<td></td>
<td>bef</td>
<td>tau</td>
<td>aft</td>
</tr>
<tr>
<td>(1)</td>
<td>(2)</td>
<td>(3)</td>
<td>(4)</td>
</tr>
<tr>
<td>Slope bef</td>
<td>slope</td>
<td>Tau</td>
<td>Tau</td>
</tr>
<tr>
<td>(1)</td>
<td>min$^{-1}$</td>
<td>min</td>
<td>min</td>
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<tr>
<td>303083</td>
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</tr>
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</tr>
<tr>
<td>SEM</td>
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<td>0.0031</td>
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</tr>
<tr>
<td></td>
<td>*</td>
<td>*</td>
<td>0.0004</td>
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</table>
Since $\dot{f}_{Ca}$ values do not show a step change, either at the beginning or after the end of hyperpolarization, the results argue against the involvement of hyperpolarization-activated channels or some type of inward rectifier channel. Another possibility would be that hyperpolarization increases the driving force for $Ca^{2+}$, which in turn would increase $Ca^{2+}$ through channels that are already open. Such an effect, however, should also be instantaneous. Furthermore, such an effect would only increase driving force by 10%, which could not explain the two-fold increase in $d/Ca/dt$.

A possible explanation could be that hyperpolarization causes the recruitment of more channels for $Ca^{2+}$ influx.

3.10 Isoproterenol Increases sarcoplasmic pH

Besides increasing the calcium content of the SR, adrenergic stimulation of a resting fiber influences the pH of the sarcoplasm as well. Fig. 30 illustrates the effect of ISO on the pH of the sarcoplasm during a 40-minute exposure. (N.B. earlier work indicated that the pK for phenol red is 0.1 – 0.4 pH units lower than the value used here of 7.7 based on in vitro measurements. The actual pH is greater by 0.1 – 0.4 pH units then indicated in Fig. 30). The preparation contained 0 Ca internal solution with 0.1 mM EGTA, the external solution was NR. Shortly after the addition of ISO myoplasmic pH increased, then after 6 minutes it decreased to a steady level, which is, on average, 0.05 pH units higher then that before ISO.
After removing ISO, there was a gradual decrease in pH, until it reached a steady level about 10 minutes later. Similar results, also with ISO and with cut fibers but 1.8 mM Ca$^{2+}$ and 20 mM EGTA in the internal solution, were obtained (unpublished).

Owen (1986) demonstrated in vascular smooth muscle that adrenergic stimulation enhances the surface Na/H exchanger via $\alpha$-adrenergic receptor pathway. She showed that ISO, though primarily a $\beta$-adrenergic receptor, stimulates $\alpha$-receptors. She found an increase in myoplasmic pH of 0.06 units, when using 10 nM ISO as agonist. This value is very close to our result in the frog skeletal muscle.

![Graph showing the effect of ISO on myoplasmic pH](image)

**Fig. 30 Effect of ISO on myoplasmic pH**

The horizontal bar and the vertical lines mark the period when 10 nM ISO was present in the external solution. Sarcoplasmic pH increased in ISO with 0.05 pH unit, and recovered to similar values after removing the drug. Internal solution contained 0 Ca and 0.1 mM EGTA, the external solution was NR. Last value of pH before ISO: 6.43, average value in ISO: 6.48, value 5 minutes after removing ISO: 6.39. Exp: 312071.
Although the literature to date lacks studies demonstrating the α-adrenergic receptor – IP₃ cascade enhancing effect on the Na/H exchanger in fast twitch skeletal muscle, it seems likely that ISO influences myoplasmic pH via this pathway.
4. DISCUSSION

4.1 Adrenergic stimulation increases $[\text{Ca}^{2+}]_{\text{SR}}$ at rest

One of the major findings of this study is that in cut fibers at rest, adrenergic stimulation increases $f_{\text{Ca}}$, indicative of an increase in the sarcoplasmic free calcium concentration ($[\text{Ca}^{2+}]_{\text{SR}}$) and thereby total SR and fiber calcium content. With normal Ringer’s external solution, ~30min adrenergic stimulation with 10 μM ISO increased $f_{\text{Ca}}$, $[\text{Ca}^{2+}]_{\text{SR}}$, and $[\text{Ca}_T]_{\text{SR}}$ (average increase in $f_{\text{Ca}}$, $[\text{Ca}^{2+}]_{\text{SR}}$, $[\text{Ca}_T]_{\text{SR}}$ were 121%, 112%, 253%, respectively). In the control experiments neither the values of $f_{\text{Ca}}$ nor those of $[\text{Ca}^{2+}]_{\text{SR}}$ and $[\text{Ca}_T]_{\text{SR}}$ showed any significant increase (Table 3.2, rows A, B, columns 2, 3 and Table 3.3, rows A, B, columns 2, 3 and 5, 6). Since the increase in $f_{\text{Ca}}$ required external calcium (Fig. 23 sagittal rows A and B), the results also indicate that ISO stimulates Ca$^{2+}$ influx across the surface/T system membrane.

These findings provide at least a partial explanation for the observations of many researchers that adrenergic stimulation increases twitch tension in frog and/or mammalian skeletal muscle (among others: Gonzalez-Serratos et al.1981, Arreola et al. 1987).

Our findings, described above, support the results of Gonzalez-Serratos and co-workers who measured an increased twitch tension for the first twitch after incubating the fiber in adrenaline without previous stimulations. They also support Rudolf et al. (2006) suggestion that stimulation with ISO results in a small increase in $[\text{Ca}^{2+}]_{\text{SR}}$. However, we are the first to show that calcium content in the SR, thereby $[\text{Ca}_T]_{\text{SR}}$ increases at rest for
adrenergic stimulation. This fact most likely leads to a greater and stronger contraction for the first stimulation measured by Gonzalez-Serratos et al. (1981).

4.2 Hyperpolarization increases Ca$^{2+}$ influx in ISO

Our major finding explained in the previous section points out that ISO stimulates Ca$^{2+}$ influx across the surface/T system membrane at rest. To evaluate whether L-type channels or some other voltage-gated Ca$^{2+}$ channels were responsible, fibers were hyperpolarized to -110 mV. Hyperpolarization to -110 mV should immediately increase the driving force for Ca$^{2+}$ by about 10%. Ca$^{2+}$ influx mediated by increased driving force would show a step change in the slope of $J_{\text{Ca}}$ that was established before hyperpolarization (Table 3.6 column 2 and Fig. 29, panel B). This slope, however, remains apparently steady for about 90 - 120s after the beginning of hyperpolarization. On the other hand, this steady slope at the beginning also indicates that voltage gated channels were not activated by ISO before hyperpolarization. If they were, we would have expected a sudden decrease of the slope of $J_{\text{Ca}}$ values, indicative of closing of these channels. The increase in rate of $J_{\text{Ca}}$ however, was coming on with a delay of $\tau = 1.6$ min (Table 3.6, column 4), until it reached its maximum final slope which is about 4-fold greater than the one established before hyperpolarization. This magnitude of increase is too great to be caused only by the increased driving force for Ca$^{2+}$ at -110 mV (Table 3.6, columns 2 and 3).
Another possibility for the increased rate of Ca$^{2+}$ entry would be the hyperpolarization-activated cyclic nucleotide-gated channels (HCN). However, the lack of step change in the rate of $\dot{f}Ca$ at the beginning of hyperpolarization period argues against their role in this process.

One interesting possible candidate is the transient receptor potential channel – Canonical (TRPC), a Ca$^{2+}$ channel without a clearly demonstrated function at this time. These channels have recently been linked to store-operated calcium entry mechanisms, but it has been suggested, that they can be activated by other unknown mechanisms (Yuan et al., 2007). In any case, the main finding in this section is that ISO stimulates some, as yet unknown, voltage-insensitive Ca$^{2+}$ channel that significantly increases the SR calcium content.

**4.3 Store-operated channels are not activated at physiologically depleted [Ca$^{2+}$]$_{SR}$**

This section considers the implications of the effects of external Ca$^{2+}$ on $\dot{f}Ca$ values in fibers with depleted [Ca$^{2+}$]$_{SR}$ before adding ISO.

In the experiments in this study, in which 20 mM EGTA was added to the internal solution, the average of initial $\dot{f}Ca$ values were $\sim$30% of those where 0.1 EGTA was present (Fig. 23 front row columns C, D vs. A, B). The average level of $\dot{f}Ca$ with 20 mM EGTA in the internal solution is the same in the presence and absence of external calcium. If surface Ca$^{2+}$ flux were significant, the values of $\dot{f}Ca$ in fibers with NR would be expected to be greater than those measured in fibers with no external Ca$^{2+}$ present. The
fact that essentially the same steady level of low \( fCa \) was measured in preparations with Ca\(^{2+}\) absent or present in the external solution indicate that store-operated Ca\(^{2+}\) entry (SOCE) is not activated at \(-30\%\) of normal resting \([Ca^{2+}]_{SR}\).

Store-operated calcium entry was demonstrated among others by Yuan et al. (2007) in transfected HEK cells, by Roos et al. (2005), in Jurkat cells, and by Stiber et al. (2008), in myotubes and adult mouse skeletal muscle cells. In the above mentioned studies, the authors depleted the SR of Ca\(^{2+}\), using thapsigargin (TG), an inhibitor of the SERCA pump. In skeletal muscle, in response to store depletion, the Ca\(^{2+}\) sensor STIM1 clusters into punctae at SR/T tubule membrane, activates ORAI1 and TRPC channels which in turn open for SOC-mediated Ca\(^{2+}\) influx (Shin and Mualem, 2008). In these preparations, SOCE was activated with near total depletion, so that it is unknown at what level of depletion gets SOCE activated.

Our study is the first to indicate that store operated mechanisms are not recruited when the calcium content changes from the physiological level as low as 30% of the physiological level. Since SOCE is not activated in this range, the question remains as to how the physiological SR calcium content is determined.

4.4 Lowered \([Ca^{2+}]_{SR}\) does not increase the rate of store filling in ISO

This section considers the implications of the effects of ISO on \( fCa \) values in fibers with normal and depleted \([Ca^{2+}]_{SR}\).
Experiments, carried out with normal \([\text{Ca}^{2+}]_{\text{SR}}\) (0.1 mM EGTA in the internal solution), showed increased rate of \(fCa\) increase in ISO when calcium was available. We made the same observation with depleted (20 mM EGTA internal) fibers. The absolute rates of increase of \(fCa\) surprisingly, were very similar in both cases (Fig. 26A, C).

It is important to note, however, that the similar rate of change of \(fCa\) in these two different preparations results in a different rate of change in \([\text{Ca}]_{\text{SR}}\). The difference is due to the non-linearity of the binding curve of Ca-Calsequestrin (CaCSQ) in this range. As Fig. 25 illustrates, the Ca-Calsequestrin binding curve is sigmoidal; its Hill coefficient is around 3, and its \(K_d = 1\) mM (Pape et al., 2007). The figure plots the binding curve of Ca-Calsequestrin the fraction of CSQ bound to \(\text{Ca}^{2+}\) \((f\text{CaCSQ}, \text{ordinate})\) vs. \([\text{Ca}^{2+}]_{\text{SR}}\) (abscissa). The range of \([\text{Ca}^{2+}]_{\text{SR}}\) at \(~30\) min. in ISO in our study is from 0.2245 mM (in depleted fibers Table 3.3, row D, column 3) to 1.2936 mM (in normal \([\text{Ca}^{2+}]_{\text{SR}}\) fibers, row A). The total length of both dotted lines in Fig 25 illustrates this entire range of \([\text{Ca}^{2+}]_{\text{SR}}\). CSQ buffering power is greater in the normal \([\text{Ca}^{2+}]_{\text{SR}}\) range, -as indicated by the steeper slope of the binding curve under the upper dotted line- than in the \([\text{Ca}^{2+}]_{\text{SR}}\) range of depleted fibers, indicated by the somewhat less steep curve at the lower dotted line. This results in a greater increase in \([\text{Ca}]_{\text{SR}}\) for the expense of the increase in \([\text{Ca}^{2+}]_{\text{SR}}\). Indeed, the average increase in \([\text{Ca}]_{\text{SR}}\) in ISO was 15.78 mM in fibers with normal \([\text{Ca}^{2+}]_{\text{SR}}\) vs. 6.07 mM in depleted fibers (Table 3.3, rows A, C, column 7).These values give an average \(d[\text{Ca}]_{\text{SR}}/dt\) of 0.5260 mM/min vs. 0.2023 mM/min in normal \([\text{Ca}^{2+}]_{\text{SR}}\) and depleted fibers, respectively. These data indicate that the \(\text{Ca}^{2+}\) influx is about 2.5 times greater in fibers with normal \([\text{Ca}^{2+}]_{\text{SR}}\).
This result is consistent with our previous finding, that store-operated mechanisms are not stimulated by ISO at this level of depletion of the SR (e.g. ~30% of normal). If ISO somehow activated the SOC-mediated Ca\(^{2+}\) entry, we would have expected a greater Ca\(^{2+}\) influx in the partially depleted fibers. This is because if SOCE was involved, one might expect the double activation to be additive i.e. ISO + store depletion.

4.5 Delay in reaching the steady rate of increase of \(\langle f \rangle_{Ca}\) in ISO

A delay in the increase of \(\langle f \rangle_{Ca}\) with an average time constant of 7.0 minutes (Fig. 27 panel A and table 3.4 part A) was seen, when ISO was added to the external solution. The time constant of the delay is consistent with the 8-minute delay in maximum force development shown by Gonzalez-Serratos et al. (1981). Interestingly, prebathing the fiber in ISO before adding Ca\(^{2+}\) eliminated the delay and caused an immediate rapid increase of \(\langle f \rangle_{Ca}\). These results together highlight that ISO does have a latent period until its effects on Ca\(^{2+}\) influx take place. It seems reasonable to hypothesize that ISO stimulates the expression of some type of channel that conducts Ca\(^{2+}\) at -90 mV.

The presence of delay in both the maximum force development and the reaching of steady rate of \(\langle f \rangle_{Ca}\) pretty much rules out the possibility that increased Ca\(^{2+}\) influx is due to activation of L-type channel via cAMP – PKA phosphorylation by the fact that this process is very rapid, takes place within a few milliseconds.

The other possibility that adrenergic stimulation may result in gene transcription and expression of a calcium-conducting cation channel in the plasmalemma seems more
likely. Microarray studies on skeletal muscle mRNA profile during epinephrine infusion showed that cAMP-dependent transcription of genes responsible for carbohydrate, lipid and protein metabolism were up-regulated by adrenergic stimulation (Arner, 2004; Viguerie et al., 2004). These include the enhanced synthesis of enzymes for glycogen mobilization, glucose metabolism, protein anabolism and inhibitory effects on proteolysis (Viguerie et al., 2004). Examining the time constant of real time gene expression in live Escherichia coli cells, Yu et al. measured a time constant of \( \tau = 7.0 \pm 2.5 \) minutes (Yu et al., 2006). The similarity of time constants shown by our results and the finding of Yu et al., suggests the possibility that adrenergic stimulation may indeed initiate gene expression.

Another possibility could be that ISO promotes the incorporation of some type of cation channel-containing vesicles into the plasmamembrane. An important biological phenomenon involving vesicle-membrane fusion is the incorporation of glucose transporters into the plasma membrane of muscle and fat cells or the fusion of ACh filled vesicles with the presynaptic membrane. Intracellular GLUT-4 traffics to the plasma membrane with a half-time of \( \sim 230 \) min, and insulin reduces the half-time to \( \sim 9 \) min (Karylowski et al., 2004). This half-time is very close to our time constant of 7.0 min. The fusion of ACh vesicles occurs on submillisecond timescales (200 \( \mu s \)) after the onset of \( \text{Ca}^{2+} \) influx (Llinás et al., 1981). SNARE-driven fusion of lipid bilayers that did not require \( \text{Ca}^{2+} \) was measured from 10 minutes half time (Weber T et al., 1998) down to a time constant of 25 ms (Liu et al., 2005). This wide range of time scale of vesicle incorporation would allow us to fit our delay towards the longer end of the range; however the absence of data would make this assumption highly speculative.
4.6 Sudden transient decrease in \( fCa \) after removal of ISO suggests change in transport of \( Ca^{2+} \) from the SR.

Another main finding of this study was the observation of rapid, transient decrease in SR calcium content while ISO was removed from the external solution. After this transient "dip", SR calcium content began increasing again with the same rate as it did during ISO (Fig. 28). These observations suggest two different effects of ISO on the fiber: one is reversible ("dip") and the other (recovery of the rate in increase) is irreversible at least on the short term. This section considers the transient decrease in \( fCa \) ("dip") after removing ISO.

The initial rate of decline of \( fCa \) after ISO removal can be described with a single exponential with a time constant of \( \tau = 0.91 \text{ min} \) (Table 3.5, column 8). From the known time constant we can easily calculate \( d[Ca_{SR}] / dt \), the rate of total calcium lost in the first minute. The calculation gives the value of 2.27 mM/min (which converts to a rate of 0.038 \( \mu \text{M/ms} \)). This rate closely corresponds to the release rate of calcium from the SR by \( \alpha \)-RyR for a depolarization to -70 mV observed previously by Pape et al., 1995 (Pape et al., 1995) in frog cut muscle fibers. Although much less than the maximum rate of \( Ca^{2+} \) release (of ~150 \( \mu \text{M/ms} \)) under the same conditions, it still corresponds to a significant rate of release. It is very important to note that any explanation for this loss of SR calcium has to involve a net efflux across the SR membrane following the removal of ISO. Because of this, it is possible to rule out a few possible explanations for the rapidly reversed component. We can approach this phenomenon from different aspects:
a.) If the removal of ISO reversed the influx pathway across the surface/T-system membrane, it would tend to stop the accumulation of Ca\(^{2+}\) in the SR but it would not cause the loss of Ca\(^{2+}\) already accumulated. Also, the observation of continuing rise of SR calcium content after the "dip" argues against the assumption that the removal of ISO closes the previously established Ca\(^{2+}\) influx pathways.

b.) Another possible explanation to consider is the hypothesis that ISO enhances Ca\(^{2+}\) leak from the SR (Reiken et al., 2003). In this case, any Ca\(^{2+}\) leak in ISO would have to be masked by the enhanced SERCA, hence the observed increase of SR calcium content during ISO. In this case, removal of ISO would reverse the leak which would cause a transient increase in SR calcium content not a decrease. The presence of the "dip" argues against the hypothesis that ISO enhances a Ca\(^{2+}\) leak from the SR.

c.) On the other hand, if ISO directly enhances SERCA by means of involvement of a stimulatory co-factor and/or phosphorylation of the pump itself, the reversal of this effect by removing ISO would expose the leak. It would do so, since prior to the removal of ISO there is a balance established with any Ca\(^{2+}\) leaked from the SR being pumped back into the SR. The reversal of the direct enhancement on SERCA would then reveal the SR Ca\(^{2+}\) leak thereby causing the "dip". In order to account for the results, this Ca\(^{2+}\) leak would also have to be reversed within several minutes following the removal of ISO. Although this explanation can not be ruled out, it is not very likely, since the "dip" indicates a very fast release, and is well described by a single exponential process.

d.) Another possibility is to evaluate the "dip" from the point of view of SERCA. Normally, the major job of SERCA is to remove Ca\(^{2+}\) from the sarcoplasm into the SR lumen. It has been shown however, that SERCA can work in reverse mode (Shannon et
al., 2000). In their study, they showed that SERCA behaves as a reversible thermodynamic enzymatic reaction where the forward rate depends on [Ca$^{2+}$]$_{myo}$, and the reverse rate on [Ca$^{2+}$]$_{SR}$. The steady state of this system occurs when the energy required to establish the [Ca$^{2+}$] gradient is balanced by the free energy, available from ATP. With the removal of ISO from the external solution, this balance shifts toward the reversal direction of the pump: [Ca$^{2+}$]$_{SR}$, and likely [ADP]$_{myo}$ and [Pi] are elevated, meantime the adrenergic enhancement of glycolytic ATP production terminates, and with this [ATP]$_{myo}$ stalls or decreases (Shannon et al. (2000). This process, however by itself would not explain the quick release of Ca$^{2+}$ from the SR. In order for this possibility to explain the rapid reversal following ISO removal, it would also require the reversal of a direct stimulation on the pump by ISO (like the ones pointed out in section c. above). This direct enhancement is necessary because the removal of ISO would not cause an immediate change in the conditions for the reversal of the pump (e.g. [Ca$^{2+}$]$_{SR}$ and [ATP]$_{myo}$).

Both explanations in section c, and d. above require in fact the hypothesis of the presence of a direct enhancing effect of ISO on the SR Ca$^{2+}$ pump; a signal pathway which is independent from the indirect enhancements due to increased [ATP]$_{myo}$ and [Ca$^{2+}$]$_{myo}$. The observed phenomenon of “dip” therefore can be considered as evidence in support of the idea that ISO directly enhances the SR Ca$^{2+}$ pump despite of the lack of apparent phosphorylation sites for PKA (Odermatt et al., 1998).
4.7 \(\text{Ca}^{2+}\) influx doesn’t change after the removal of ISO

Fig. 28 shows that after the lowest point of the “dip” \(\dot{\text{f}}\text{Ca}\) values begin to rise, until they reach a steady level. The slope of line fit of the steady level after the removal of ISO is not significantly different from the one established in ISO (Table 3.5, columns 2, 3). This finding indicates that removal of ISO does not change the \(\text{Ca}^{2+}\) influx present before removing ISO. It seems reasonable to conclude that the similar rate of \(\text{Ca}^{2+}\) influx is due to expressed \(\text{Ca}^{2+}\) channels in response to ISO stimulation or some type of very slow up-regulation of \(\text{Ca}^{2+}\) channels, already present when ISO was added.

4.8 \(\text{Ca}^{2+}\) leak from the SR in ISO is unlikely

As discussed above, the “dip” argues against the enhancement of a \(\text{Ca}^{2+}\) leak by ISO. This section gives another reason to question this hypothesis.

When we exposed the fiber containing 20 mM EGTA and 0 \(\text{Ca}^{2+}\) in the internal solution to ISO there was no significant change in the rate of change of \(\dot{\text{f}}\text{Ca}\) values in ISO (Fig. 22 panel D, and Fig. 26, D). If there was a leak, we would expect to see a transient decrease in \(\dot{\text{f}}\text{Ca}\). This would be expected, since the 20 mM EGTA would buffer the leaked \(\text{Ca}^{2+}\), and would compete with SERCA. This process would delay the pumping back of leaked \(\text{Ca}^{2+}\) in to the SR. This observation provides direct evidence against any \(\text{Ca}^{2+}\) leak from the SR occurring during ISO exposure; also it argues against the
hypothesis made by Marks and co-workers (Reiken et al., 2003) that adrenergic stimulation causes the RyR channel to be leaky.

4.9 Enhancement of α-adrenergic pathway can explain myoplasmic pH change in ISO

Fig. 30 illustrates the change of myoplasmic pH during ISO. After an initial transient increase pH values gradually decreased to a steady value of 0.05 pH unit above the initial value, and remained at this elevated level until ISO was washed out. After washing out ISO, myoplasmic pH recovered to the value before adding ISO.

Regulation of myoplasmic pH depends on continuous activity of membrane transport systems that mediate an outflux of $\text{H}^+$ or influx of bicarbonate, so that proton generation is counterbalanced by proton efflux (JUEL, 1998). The transporters involved in muscle pH regulation at rest are the $\text{Na}^+/\text{H}^+$ exchange system (Moore, 1981; Owen, 1986), as well as the $\text{Na}^+$-dependent (NBCe), and $\text{Na}^+$-independent (Cl/HCO$^3$-X) Cl - bicarbonate transport systems (Wang et al., 2000; J. M. Kristensen, 2004). Proton-linked monocarboxylate transporters (MCT) are also present in skeletal muscle. These proteins are responsible for efflux of glycolytically derived lactic acid. In fast twitch skeletal muscle, MTC3 appears to be the major isoform (Wilson et al., 1998).

Lactate/H$^+$ transporters (MCT) appear to be the major acid extruding system associated with intense exercise (JUEL, 1998), while the $\text{Na}^+/\text{H}^+$ exchanger and the
Na⁺/HCO₃⁻ co-transporter (NBC) seem to be active at resting myoplasmic pH levels in skeletal muscle (JUEL, 1998; J. M. Kristensen, 2004).

Since our experiments revealed a myoplasmic pH rise during adrenergic stimulation, the major question is, whether these proteins responsible for pH, can be affected by isoproterenol?

Owen (1986) proposed that isoproterenol interacts with α-adrenergic receptor to cause IP₃ to rise, which in turn, activates Na/H exchange. This effect could be blocked by the α-antagonist phenolamine, but not by the β-antagonist propranolol. In her studies, she found that 10 μM ISO increased basal pH by 0.06 pH unit (Owen, 1986) which is very close to the pH increase of 0.05 in ISO, revealed in our experiment. Our results are very similar to earlier, unpublished results with 20 mM EGTA in the internal solution with frog cut fibers from our laboratory (Pirouzi, 1999). His results indicated a 0.05 pH increase, reached after a 4-minute incubation in 10 μM ISO. After removing ISO from the external solution, myoplasmic pH also recovered completely suggesting that isoproterenol is the most likely cause of pH rise.

There is another protein which is responsible for the acid-base homeostasis of skeletal muscle at rest, the Na⁺/HCO₃⁻ co-transporter (NBCe). Kristensen et al., 2004 (J. M. Kristensen, 2004) revealed, that approximately half of the pH recovery is mediated by bicarbonate-dependent transport, by NBCs. They localized two isoforms of the protein in sarcolemmal membranes and suggested that they are present in some internal membranes, probably the T-tubules. The location and function of NBCs would make them another ideal candidate besides the Na/H exchanger for increasing myoplasmic pH if they could be enhanced by adrenergic stimulation. Indeed, Hayashi, 2001 (Hayashi, 2001) reported
that NBCs, found in kidney β-intercalated cells are controlled by isoproterenol via β₁-receptor-cAMP-PKA pathways, but this function has not been verified in skeletal muscle.

Since bicarbonate is not present in our fibers, our results suggest that the increase in myoplasmic pH caused by isoproterenol is likely due to enhanced Na/H effect stimulated via PLC-IP₃ pathway.
5. CONCLUSION & PERSPECTIVES

The data presented in this thesis show new findings on the effects of isoproterenol on $[\text{Ca}^{2+}]_{\text{SR}}$ and $[\text{Ca}_{\text{t}}]_{\text{SR}}$ at rest using TMX, a membrane permeable low affinity Ca$^{2+}$-indicator. We monitored the fraction of Ca$^{2+}$-bound TMX ($\gamma\text{Ca}$), a direct indicator of free calcium content in the SR ($([\text{Ca}^{2+}]_{\text{SR}})$.

1. We established that adrenergic stimulation increases Ca$^{2+}$ influx across the sarcolemma in resting fiber. This increased Ca$^{2+}$ influx is non-reversible within the time period of tens of minutes after removing adrenergic stimulation.

2. The isoproterenol effect occurs with a delay with an exponential time constant of $\tau = 7.0 \text{ min}$ until the rate of Ca$^{2+}$ influx reaches a steady level. On the other hand, Ca$^{2+}$ influx in fibers with ISO in the external solution instantaneously increased upon the addition of external Ca$^{2+}$. Furthermore, after removing ISO, following a transient "dip" the rate of Ca$^{2+}$ influx recovered, and remained constant at the same rate that was established in ISO. These observations suggest that adrenergic stimulation promotes the expression of some type of channel that conducts Ca$^{2+}$ at rest and these channels remain active after the removal of adrenergic stimulation.

3. We showed that the rate of increase of $\gamma\text{Ca}$ established in ISO increases after some delay following hyperpolarization. Because of the delay, the increase can not be
explained simply by the increased driving force for Ca\(^{2+}\) or by inward rectification of a Ca\(^{2+}\) conducting channel or by hyperpolarization activated channels. These data suggest therefore that ISO stimulates some, as yet unknown voltage-insensitive Ca\(^{2+}\) channel that significantly increases the SR calcium content. Putative candidate may be related to the TRP super family of canonic channels which would be activated at rest.

4. We demonstrated that Ca\(^{2+}\) influx across the sarcolemma hence \([\text{Ca}^{2+}]_{\text{SR}}\) is not enhanced in fibers with lowered \([\text{Ca}^{2+}]_{\text{myo}}\) to 30\% of its normal level. This finding argues against the participation of store-operated mechanisms at this level of depletion.

5. The steady level of \(\text{Ca}^{2+}\) in ISO in fibers containing no Ca\(^{2+}\) in the external solution and 20 mM EGTA in the internal solution argues against the hypothesis of Marks and colleagues that adrenergic stimulation causes Ca\(^{2+}\) leak from the SR. The phenomenon of the "dip", the transient decrease in \([\text{Ca}^{2+}]_{\text{SR}}\) after the removal of ISO, also argues against the hypothesis that ISO enhanced a Ca\(^{2+}\)-leak from the SR during adrenergic stimulation.

6. The occurrence of the "dip", -which is equivalent with a Ca\(^{2+}\) release from the SR for a depolarization to -70 mV- its magnitude of Ca\(^{2+}\) release, its single exponential time course, and its transient nature suggest that ISO directly enhances the SR Ca\(^{2+}\) pump and the removal of ISO brings this enhancement to an end. This enhancement may occur by a direct enhancement of a co-stimulatory factor and/or phosphorylation of the pump itself if possible. After the removal of ISO, with no further adrenergic stimulation the pump quickly turns into "reverse mode" due to the high \([\text{Ca}^{2+}]_{\text{SR}}\).
7. Our results show an increase in pH by ISO. This pH increase supports the possibility of the involvement of α-adrenergic pathways and its downstream events. Although isoproterenol is primarily a β-adrenergic agonist, its α-adrenergic influence—though in much less extent than that of β—has been previously shown.

In summary, ISO increases Ca$^{2+}$ influx across the surface/T-system membrane which in turn increases SR calcium content ([Ca]$^{2+}$) and with this, [Ca$^{2+}$]$_{SR}$. Increased [Ca$^{2+}$]$_{SR}$ leads to a greater Ca$^{2+}$ release from the SR which causes a greater tension during muscle contraction.

Our results may give physiological importance to the fact that athletes, directly before their race, relax in a motionless position. It is possible that this concentration before the race not only prepares them mentally but physically as well through the effects of elevated adrenaline levels on their muscles shown in this study.

On the other hand, bedridden patients might benefit from adrenergic stimulation administered directly before their regular physical therapy exercise. The enhancing effects of adrenaline on following muscle contractions could help better maintain—or slower the rate of atrophy—of muscles.
6. APPENDIX - A

The purpose of this appendix is to describe the approach used to calculate the best fit, using the least sum of squares differences method for the "dip", the transient decrease of fCa values after the removal of ISO. This phenomenon is illustrated in Fig 27. After removing ISO, fCa values decreased quickly, reaching their lowest value, then recovered, reaching a steady, positive slope. The shape of this transient decrease could be best described by an exponential function with a sloping line (y_{fit}) described by eqn. 3.14:

\[ y_{fit} = C \left( 1 - e^{-\frac{t'}{\tau}} \right) + mt \]  \hspace{1cm} (A-1)

where \( C \) is the magnitude of exponential, \( m \) is the sloping line, reached at \( t = \infty \), and

\[ t = t' - t_{REF}, \] \hspace{1cm} (A-2)

where \( t' \) is time after the addition of DYE, and \( t_{REF} \) is the time when \( fCa \) begun to drop, so \( t = 0 \) at the beginning of the fit.

The value of \( fCa \) at \( t = 0 \) was set to 0, and denoted as \( y \):

\[ y = fCa - fCa(t_{REF}), \] \hspace{1cm} (A-3)
The least sum of squares fit between the function in eqn. A-1 and y values:

$$SSQ = \sum_{i=1}^{N} (y_i - y_{fit})^2$$  \hspace{1cm} (A-4)$$

where $y_i$ are values of $fCa$ of time $(t_i)$ below, in eqn. A-5, after the removal of ISO.

By substituting the function of $y_{fit}$ into eqn. A-4:

$$SSQ = \sum_{i=1}^{N} \left[ y_i - \left( C \left( 1 - e^{-t_i} \right) + mt_i \right) \right]^2$$  \hspace{1cm} (A-5)$$

We obtain the best fit, if the partial derivative of SSQ in respect to $m$ and $C$ are 0. By solving eqn. A-5, first in respect to $m$:

$$\frac{\partial SSQ}{\partial m} = -2\sum_{i=1}^{N} \left[ y_i - \left( C \left( 1 - e^{-t_i} \right) + mt_i \right) \right] t_i = 0,$$  \hspace{1cm} (A-6)$$

rearranging,

$$\sum y_i t_i - C\sum t_i + C\sum t_i e^{-t_i} - m\sum t_i^2 = 0,$$  \hspace{1cm} (A-7)$$
and solving for $m$:

$$m = - \frac{C \sum t_i - C \sum t_i e^{t_i} + \sum y_i t_i}{\sum t_i^2}.$$  \hfill (A-8)

Solving the partial derivative of SSQ in respect to $C$:

$$\frac{\partial \text{SSQ}}{\partial C} = -2 \sum y_i - \left( C \left( 1 - e^{t_i} \right) + mt_i \right) \left( 1 - e^{t_i} \right) = 0$$ \hfill (A-9)

rearranging:

$$\sum y_i - CN + C \sum e^{t_i} - m \sum t_i = \sum y_i e^{t_i} + C \sum e^{t_i} - C \sum e^{t_i} + m \sum t_i e^{t_i} = 0.$$ \hfill (A-10)

Substituting in eqn. A-8, and combining $C \sum e^{t_i}$ terms:

$$\sum y_i - \sum y_i e^{t_i} + 2C \sum e^{t_i} - C \sum e^{t_i} - CN +$$

$$+ \frac{C \left( \sum t_i - \sum t_i e^{t_i} \right) - \sum y_i t_i}{\sum t_i^2} \left( \sum t_i e^{t_i} - \sum t_i \right) = 0,$$ \hfill (A-11)
and finally solving for $C$:

$$C = \frac{\sum t_i \left( \sum y_i e^{\frac{-2t_i}{T}} - \sum y_i \right) - \sum y_i t_i \left( \sum t_i e^{\frac{-2t_i}{T}} - \sum t_i \right)}{\sum t_i^2 \left( 2\sum e^{\frac{-2t_i}{T}} - \sum e^{\frac{-2t_i}{T}} - N \right) + \left( \sum t_i - \sum t_i e^{\frac{-2t_i}{T}} \right)^2}.$$  \hspace{1cm} (A-12)

The function of the fit, $y_{FIT}$ (A-1) with the solutions for variables $m$ (A-8) and $C$ (A-12) were entered in an excel worksheet. The best fit between values of $y_i$ and $y_{FIT}$ were obtained by getting the least sum of squares differences between the two sets of values by changing the user-adjustable parameter, $T$:

$$SSQ_{y,y_{FIT}} = \sum (y_i - y_{FIT})^2 \hspace{1cm} (A13)$$
APPENDIX - B

The aim of this appendix is to describe the approach of calculating \([Ca^{2+}]_{SR}\) from the measured \(f_{Ca}\) values.

Since calcium bound TMX is found explicitly in the SR, and taking the distribution of TMX forms in the fiber into account, the following formula describes \(f_{Ca}\):

\[
f_{Ca} = \frac{V_{SR}[CaTMX]_{SR}}{V_{SR}([CaTMX]_{SR}+[TMX]_{SR})+V_{myo}[TMX]_{myo}}
\]  

(B1)

where \(V\) is volume and the subscripts \(SR\) and \(myo\) refer to sarcoplasm and myoplasm, respectively. This value, however, underestimates SR free calcium, because it contains the myoplasmic component of TMX ([TMX]_{myo}). To calculate the fraction of calcium bound TMX in the SR (\(f_{Ca,SR}\)) we have to omit TMX, present in the myoplasm:

\[
f_{Ca,SR} = \frac{V_{SR}[CaTMX]_{SR}}{V_{SR}([CaTMX]_{SR}+[TMX]_{SR})}.
\]  

(B2)

Combining eqns (B1) and (B2) and expressing the reciprocal of \(f_{Ca}\) yields:

\[
\frac{1}{f_{Ca}} = \frac{1}{f_{Ca,SR}} + \frac{V_{myo}[TMX]_{myo}}{V_{SR}[CaTMX]_{SR}}.
\]  

(B3)
The second part of the right side of eqn (B3) can be broken down into two parts knowing that

\[ [TMX]_{SR} = \rho [TMX]_{myo}, \]  

(B4)

where \( \rho \) is a constant that explains the difference in \([TMX]_{myo}\) and \([TMX]_{SR}\) based on the pH gradient across the SR membrane. Substituting the ratios of TMX forms present in the SR in (B3) with their fractional ratios and dividing with \( \rho \) yields:

\[
\frac{V_{myo}}{V_{SR}} \cdot \frac{[TMX]_{myo}}{[CMX]_{SR}} = \frac{V_{myo}}{\rho V_{SR}} \cdot \frac{1 - fCa_{SR}}{fCa_{SR}},
\]  

(B5)

and substituting eqn (B5) into the second expression of the right side of eqn (B3) gives the final solution:

\[
\frac{1}{fCa} = \frac{1}{fCa_{SR}} + \frac{V_{myo}}{\rho V_{SR}} \cdot \frac{1 - fCa_{SR}}{fCa_{SR}}.
\]  

(B6)

The volume ratio \( (V_{myo}/V_{SR}) \) is a constant, as well as \( \rho \), therefore eqn (B6) describes the function of \( fCa_{SR} \) vs. \( fCa \). Following these calculations with the monitored \( fCa \) values, \([Ca^{2+}]_{SR}\) can be calculated with eqn 2.21:
\[ [\text{Ca}^{2+}]_{\text{SR}} = \frac{f\text{Ca}_{\text{SR}} \cdot K_d}{1 - f\text{Ca}_{\text{SR}}}. \]  

It is important to note, however, that in case the turnover rate of the SERCA changes, it should change the constant \( \rho \) (which is obtained from the pH gradient across the SR membrane eqn. 3.5), which in turn will change slightly the relationship between \( f\text{Ca} \) and \( f\text{Ca}_{\text{SR}} \) and therefore the value of \([\text{Ca}^{2+}]_{\text{SR}}\).
This section describes how information about the concentrations of Ca-bound and Ca-free TMX ([CaTMX] and [TMX], respectively) and related information is derived from the measurement-derived \( A_{TMX} \) signals. Since there are two dye species, TMX and CaTMX, their concentrations can be determined from the dye-related absorbances at two wavelengths. This section describes how [TMX] and [CaTMX] can be obtained utilizing the information from two or more wavelengths using the least-squares method. The measurement-derived absorbance at wavelength \( \lambda_n \) is denoted \( A_{dye}(\lambda_n) \) where \( n \) can vary from 1 to the number of wavelengths, \( N \). The least-squares best fit absorbance at wavelength \( \lambda_n \) is given by

\[
A_{fit}(\lambda_n) = \varepsilon_{TMX}(\lambda_n) \cdot [TMX] \cdot l + \varepsilon_{CaTMX}(\lambda_n) \cdot [CaTMX] \cdot l
\]  

(C1)

where \( l \) is the optical path length and \( \varepsilon_{TMX}(\lambda_n) \) and \( \varepsilon_{CaTMX}(\lambda_n) \) are the extinction coefficients at wavelength \( \lambda_n \). This is simply the sum of the absorbances from each of the dye species using Beer's Law and the best-fit values of [TMX] and [CaTMX]. The sum of squares (SSQs) is given by

\[
SSQs = \sum_{n=1}^{N} (A_{dye}(\lambda_n) - A_{fit}(\lambda_n))^2
\]  

(C2)

Substituting equation (C1) into (C2), taking the partial derivatives of SSQs with respect to [TMX] and [CaTMX] and solving for [TMX] and [CaTMX] from the two partial derivatives gives that
\[ [\text{TMX}] = \frac{\sum_{\nu} \varepsilon_{\text{CatMx}}^2 \cdot \sum \Delta_{\nu} \varepsilon_{\text{TMx}} - \sum \varepsilon_{\text{TMx}} \varepsilon_{\text{CatMx}} \cdot \sum \Delta_{\nu} \varepsilon_{\text{CatMx}}}{\sum \varepsilon_{\text{TMx}}^2 \sum \varepsilon_{\text{CatMx}}^2 - \left( \sum \varepsilon_{\text{TMx}} \varepsilon_{\text{CatMx}} \right)^2} \] (C3)

and

\[ [\text{CatMx}] = \frac{\sum \varepsilon_{\text{TMx}}^2 \cdot \sum \Delta_{\nu} \varepsilon_{\text{CatMx}} - \sum \varepsilon_{\text{TMx}} \varepsilon_{\text{CatMx}} \cdot \sum \Delta_{\nu} \varepsilon_{\text{CatMx}}}{\sum \varepsilon_{\text{TMx}}^2 \sum \varepsilon_{\text{CatMx}}^2 - \left( \sum \varepsilon_{\text{TMx}} \varepsilon_{\text{CatMx}} \right)^2} \] (C4)

(N.B. for brevity, the wavelength index, \( n \), for the summations and the function dependencies on \( \lambda_n \) are omitted.)
APPENDIX - D

This section describes the steps used for calculating the time constant of the delay depicted in Fig. 27 A.

The function that describes the change of $fCa$ in ISO, is based on an exponential function and the rate of changes of $fCa$ before ISO and at infinite time, where the rate of change becomes steady:

\[
\frac{dfCa}{dt} = \left( \frac{dfCa}{dt} \bigg|_{\infty} - \frac{dfCa}{dt} \bigg|_{0} \right) \left( 1 - e^{-\frac{t}{\tau}} \right) + \frac{dfCa}{dt} \bigg|_{0}
\]

where $t$ is the time of addition of ISO,

and $\left. \frac{dfCa}{dt} \right|_{0}$ is the slope of $fCa$ before ISO, denoted $m_o$,

and $\left. \frac{dfCa}{dt} \right|_{\infty}$ is the steady slope of $fCa$, reached where $t >> \tau$, denoted $m_\infty$.

After substituting $m_0$ and $m_\infty$ in to eqn E1, it simplifies:

\[
\frac{dfCa}{dt} = (m_\infty - m_0) \left( 1 - e^{-\frac{t}{\tau}} \right) + m_0.
\]

Integration gives:

\[
fCa(t) = fCa(0) - (m_\infty - m_0) \left( 1 - e^{-\frac{t}{\tau}} \right) + m_\infty t
\]

where $fCa(0)$ is the value of $fCa$ at the time of addition of ISO
Define the function of \( f^{\text{Ca}_{\text{FIT}}} \), the line fit on \( f^{\text{Ca}} \) values when they reached a steady slope as follows:

\[
f^{\text{Ca}_{\text{FIT}}} = m_{-} t + f^{\text{Ca}_{\text{FIT}}}(0),
\]

where "\( f^{\text{Ca}_{\text{FIT}}}(0) \)" is the value of the fit at the addition of ISO, depicted in Fig. 27 A.

In the case, when \( t \gg 0 \), \( f^{\text{Ca}_{\text{FIT}}} = f^{\text{Ca}} \), so eqn E3 and E4 gives:

\[
m_{-} t + f^{\text{Ca}_{\text{FIT}}}(0) = f^{\text{Ca}}(0) + m_{-} t - (m_{-} - m_{0}) r,
\]

and finally by rearranging, we can express the time constant:

\[
\tau = \frac{f^{\text{Ca}}(0) - f^{\text{Ca}_{\text{FIT}}}(0)}{m_{-} - m_{0}}.
\]
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