Electrophysiology and Dye-Coupling Are Sexually Dimorphic Characteristics of Individual Laryngeal Muscle Fibers in *Xenopus laevis*

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Sex differences at the laryngeal neuromuscular junction of *Xenopus laevis* were examined by recording intracellularly from muscle fibers in response to nerve stimulation. Male laryngeal muscle contains 2 physiologically distinct fiber types. Type I fibers generate postsynaptic potentials in response to low-magnitude stimulus pulses and action potentials in response to higher-magnitude stimulus pulses. Type II muscle fibers require repetitive stimulation for action potential production, probably because of facilitation. Subthreshold events in type I and II fibers suggest that these neuromuscular synapses have low safety factor junctions. Female laryngeal muscle contains one fiber type (III), which is physiologically distinct from those found in the male. Type III fibers produce an action potential in response to a single-stimulus pulse of suprathreshold voltage delivered to the laryngeal nerve; subthreshold events were not observed. Lontophoretic injection of Lucifer yellow into a single female muscle fiber resulted in as many as 43 labeled fibers. In males, only one fiber was labeled. Dye-coupling was not observed in adult females treated with the androgenic steroid hormone, testosterone. We have previously reported that laryngeal muscle fibers are recruited throughout a stimulus train presented to the laryngeal nerve in males, but are not recruited in females (Tobias and Kelley, 1987). Sex differences in the frequency of electrophysiological fiber types described here may account for sex differences in fiber recruitment. Synchronous activity of dye-coupled fibers may increase the effectiveness of muscle contraction in females.

The clawed frog, *Xenopus laevis*, relies on vocal behavior to advertise reproductive state. Vocalizations are sexually dimorphic in rate, temporal pattern, and function (Wetzel and Kelley, 1983). The male-specific mate call—a rapid trill with alternating fast and slow click rates—attracts and excites females (Picker, 1983). The female-typical call, ticking—a slow monotonous trill—is used to terminate male clasping attempts (Weintraub et al., 1985). Sounds are produced when laryngeal muscles contract, pulling open 2 apposed discs of artenoid cartilage (Yager, 1982). These distinctive vocalizations differ in requirements for laryngeal muscle activity. Mate calling requires laryngeal muscles to contract rapidly (71 Hz) for long periods (Wetzel and Kelley, 1983). Ticking requires laryngeal muscles to contract more slowly (6 Hz) for short periods (Hannigan and Kelley, 1986).

Sex differences in behavior can be due to male/female dimorphisms in the CNS and/or the periphery. Laryngeal motor neurons (Simpson et al., 1986; Kelley et al., 1988), CNS pathways (Wetzel et al., 1985), and laryngeal muscle (Sassoon and Kelley, 1986; Sassoon et al., 1987; Segal et al., 1987) are all sexually dimorphic candidates for producing sex differences in vocal behavior. Recent studies in the isolated larynx, examining muscle activity in response to nerve stimulation (Tobias and Kelley, 1987), reveal that at least 2 properties of call production are regulated in the periphery: the rate of muscle contraction, which controls the rate of click production, and the strength of muscle contraction, which contributes to click amplitude. The rate at which discrete tension transients (muscle contractions that increase and decrease to resting level) are produced by laryngeal muscle in response to nerve stimulation is considerably faster in males than in females. Histochromical characteristics of laryngeal muscle are consistent with the dimorphism in contraction rate. Male laryngeal muscle is composed entirely of fast-twitch, highly oxidative fibers, while female laryngeal muscle is composed primarily of slow, moderately oxidative fibers (Sassoon et al., 1987). The size of muscle tension transients is controlled by the number of muscle fibers contracting simultaneously, a property reflected in the amplitude of muscle compound action potentials. In male laryngeal muscle, nerve-evoked compound action potential amplitudes increase dramatically throughout the entire stimulus train. In female laryngeal muscle, compound action potential amplitudes increase significantly less and only at the beginning of the stimulus train. The magnitude of potentiation is directly proportional to stimulus frequency in males and inversely proportional in females. Since each compound action potential is the summed response of all muscle fibers responding to a given stimulus pulse, we suggest that only laryngeal muscle fibers recruited throughout the stimulus train. Possible sites for regulation of muscle fiber recruitment in the isolated larynx are the muscle fiber membrane or laryngeal motor neuron terminals.

One goal of the present study was to examine possible sex differences in the electrophysiological properties of laryngeal muscle fibers. These experiments extend our previous work (Tobias and Kelley, 1987) to the cellular level. In addition, we wished to examine mechanisms underlying sex differences in fiber recruitment. We report here that laryngeal muscle is composed of 3 electrophysiologically distinct fiber types, which can be classified as male- or female-like on the basis of their relative abundance in the 2 sexes. The frequency and characteristics of

Received July 8, 1987; revised Dec. 4, 1987; accepted Dec. 10, 1987.

This work was supported by NIH Grants NS 23684 and NS 07685.

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each fiber type within a sex are consistent with observed sex differences in fiber recruitment. Androgen treatment of adult females does not significantly increase the proportion of male-like fibers. The relative contributions of pre- and postsynaptic mechanisms to observed electrophysiological fiber types are discussed.

A second goal of this study was to examine the relation between muscle fiber types as characterized electrophysiologically (this study) and histochemically (Sassoon et al., 1987). Studies of ATPase and SDHase expression in laryngeal muscle show that fiber diameter is correlated with twitch type (Sassoon et al., 1987). Thus, to compare electrophysiological fiber type with diameter, we injected individual fibers with the low-molecular-weight dye Lucifer yellow (Stewart, 1978). We have found that, in females, filling a single fiber with dye can result in the labeling of many fibers of different diameters. In males, dye filling a single fiber always results in only one labeled fiber. The presence of dye-coupling in laryngeal muscle appears to be androgen-regulated, since coupled fibers have not been observed in male or in testosterone-treated female laryngeal muscle. We hypothesize that adult female laryngeal muscle fibers form functional syncytia that can be disrupted by the androgenic steroid hormone testosterone.

Materials and Methods

Adult *Xenopus laevis* were obtained from *Xenopus* I (Ann Arbor, MI) or Nasco (Fort Atkinson, WI). Animals were maintained on a 12:12 hr light:dark cycle in plastic tanks, and fed frog brittle every other day. Some adult females were briefly anesthetized with cold and received a 16-20 mg implant of testosterone propionate (Sigma) into the dorsal lymph sac for 1 month. Females in this study were not ovariectomized because a previous study had shown no differences in laryngeal muscle physiology (compound action potential potentiation and contraction rates) between intact, hormone-implanted and ovariectomized, hormone-implanted females (Tobias and Kelley, 1987). Prior to dissection, all frogs were anesthetized by immersion in ethyl m-aminobenzoate methanesulfonic acid (0.1%; Aldrich). The larynx was excised as previously described (Tobias and Kelley, 1987), pinned dorsal side up in a wax-coated petri dish, and bathed in glucose-supplemented amphibian Ringer solution (NaCl, 0.1 m; KCl, 2 mm; CaCl2, 2.5 mm; MgCl2, 3.0 mm; HEPES, 4 mm; glucose, 27.7 mm). Laryngeal muscle is capable of strong contractions for several hours under these conditions (Tobias and Kelley, 1987). Individual muscle fibers were impaled with glass microelectrodes filled with either potassium acetate (2.5 M) or Lucifer yellow (4% in 0.1 M LiCl). Electrode resistance ranged from 25 to 45 MΩ. Recordings were amplified (Getting, model 5A) and photographed directly from a storage oscilloscope (Tektronix S113). Only fibers with resting potentials greater than 80 mV were examined. The laryngeal nerve was stimulated via a suction electrode (WPI 302-T). The stimulus voltage was gradually increased until a muscle fiber response was elicited. Stimulus trains were 200 msec in duration and varied in frequency from 25 to 66 Hz. All stimulus trials were separated by 10 sec, to avoid facilitation or summation at the postsynaptic membrane.

Lucifer yellow was iontophoretically injected (5-10 nA, 500 msec/sec) into single fibers. Larynges containing dye-filled cells were placed in fresh 4% paraformaldehyde (0.1 M phosphate buffer) for 3 hr, followed by 20% sucrose/paraformaldehyde overnight. Tissue was cryostat-sectioned (25 μm), dehydrated, cleared in xylene, and mounted on a non-fluorescent medium (Lustrex; Monsanto). Sections were examined with a Zeiss fluorescence microscope. The significance of the differences in frequency of fiber types encountered was evaluated using the chi-square test (Siegel, 1956). The sex differences in the frequency of dye-coupled fibers were evaluated using the Fisher exact probability test (Siegel, 1956).

Results

Physiological sex differences in laryngeal muscle fibers

Males. Male laryngeal muscle contains 2 electrophysiologically identifiable fiber types. Type I fibers generate action potentials in response to single-stimulus pulses delivered to the laryngeal nerve, while type II fibers generate action potentials only in response to stimulus trains. Response of a type I fiber are illustrated in Figure 1. A low-voltage stimulus pulse presented to the laryngeal nerve results in a single subthreshold event in the muscle fiber (Fig. 1A). As the voltage is increased, a postsynaptic potential (PSP) of similar amplitude is produced. Still higher-voltage stimulation of the laryngeal nerve results in production of an action potential. The amplitude of the response, PSP, or action potential, is solely dependent on stimulus strength, regardless of the order in which the stimulus pulses are presented. Stimulus trains at voltages subthreshold for action potential generation initially produce a PSP, followed by action potentials (Fig. 1B). Action potentials are then generated in response to each stimulus pulse within the train.
Figure 2. Responses of 2 type II fibers to nerve stimulation in male larynx. A. An action potential is not produced in response to single-stimulus pulses at 2 stimulus strengths (top). This fiber is relatively frequency-independent in that an action potential is produced after the fourth subthreshold event in response to a range of stimulus frequencies (lower 3 panels). Note that failures can occur after the production of action potentials (bottom). B. As with the fiber shown in A, no action potential is produced in response to a single-stimulus pulse (top). As the stimulus frequency is increased, the number of action potentials per unit time increases, and the latency to the first action potential decreases. Note that failures to generate action potentials occur within the train even after an action potential has been generated (36 and 20 ms). Calibration bars (A, B): 10 mV (vertical), 5 msec (horizontal, top), 20 msec (lower 3 panels). ms, milliseconds.

Type II fibers produce only subthreshold events in response to single-stimulus pulses delivered to the laryngeal nerve (Fig. 2). The amplitude of the PSP is the same regardless of stimulus strength. Stimulus pulses were presented at the same magnitude as, or greater than, those required to elicit an action potential in type I fibers. Type II fibers vary in the extent to which they are frequency-dependent; 2 examples are shown in Figure 2. In the frequency-sensitive fiber, action potentials are produced after fewer subthreshold events as the interpulse interval is decreased (Fig. 2B). In the frequency-insensitive fiber, action potentials are produced after the same number of subthreshold events over a similar range of stimulus frequencies (Fig. 2A). PSP amplitude tends to increase with succeeding stimuli within a train, indicating facilitation. Type II fibers do not generate an action potential in response to every stimulus pulse within a train, even after the initial action potential is produced.

Females. Female laryngeal muscle contains predominantly one fiber type (type III), which is electrophysiologically distinguishable from those types found in male laryngeal muscle. Type III fibers produce an action potential in response to single-stimulus pulses of suprathreshold voltage delivered to the laryngeal nerve (Fig. 3A). Subthreshold events were not observed in the 38 fibers examined. Stimulus trains elicit an action potential in response to each pulse within the train (Fig. 3B). We have ob-
Sex differences in dye-coupling of laryngeal muscle fibers

In some preparations, a Lucifer yellow-filled electrode was used to iontophoretically inject dye into a muscle fiber following electrophysiological characterization. Dye injection into male laryngeal muscle fibers resulted in only one labeled cell in all fibers examined (Fig. 4A, Table 1). In contrast, dye injection into a female laryngeal muscle fiber sometimes (in 8 out of 20 cases) resulted in multiple labeling (Fig. 4B, Table 1). Both type I and type III fibers were dye-coupled. Both small and large fibers were dye-coupled. Thus, it is likely that fibers of various physiological types are coupled together. The number of labeled cells in the dye-coupled preparations varied from 2 to 43. Laryngeal muscle fibers are organized in an anterior-to-posterior direction. Dye-coupled fibers are organized in rows; any one fiber resulted in multiple-cell labeling, the diameter of the impaled cell could not be determined. Fiber diameters of 3 single-labeled fibers were compared to those of other fibers in the same muscle. In 2 preparations, the dye-filled fiber was similar in diameter to the largest fibers in the same muscle. In one preparation, the dye-filled fiber was similar in diameter to medium-sized fibers in the same muscle. Thus, large- and medium-sized fibers were impaled with Lucifer yellow-filled electrodes.

Effects of testosterone on female laryngeal muscle fibers

Testosterone treatment of adult females results in partial masculinization of calling behavior (Hannigan and Kelley, 1986), of muscle fiber size and histochemical fiber types (Sassoon et al., 1987), and of laryngeal muscle tension characteristics and compound action potential amplitude (Tobias and Kelley, 1987). An increase in the percentage of male-typical physiological fiber types could help to explain the masculinization of compound action potential amplitude potentiation (Tobias and Kelley, 1987). The frequency of male-typical muscle fibers (types I and II) is not significantly different in control and testosterone-treated females. Testosterone-treated females continue to exhibit a predominance of type III fibers.

Lucifer yellow injection into laryngeal muscle fibers from testosterone-treated females resulted in only one labeled cell per

Table 1. Incidence of physiological fiber type and dye-coupling in male, female, and testosterone-treated female frogs

| Type | Male (%) | T-Female (%) | Female (%) |
|------|----------|--------------|------------|
| I    | 2 (25.0) | 2 (12.5) | 2 (4) |
| II   | 31 (70.5) | 2 (12.5) | 7 (15) |
| III  | 2 (4.5) | 12 (75.0) | 38 (81) |
| Total | 44 | 16 | 47 |

| Dye-coupled/[total] | Male | T-Female | Female |
|---------------------|------|----------|--------|
| 0 [15]              | 0 [8] | 8 [20]   |

* Total number of fibers examined in 20 males, 8 testosterone-treated females, and 21 females.

* Instances of dye-coupled fibers [total number of fibers injected]. This experiment was performed in 9 males, 5 testosterone-treated females, and 9 females.

* The frequency of dye-coupling in females is significantly greater than that in males (Fisher exact probability test; p = 0.017). In 2 of the 8 cases in which coupling was observed, only 2 fibers were coupled. These cases were not included in the statistical analysis.

Figure 3. Response of a type III fiber to nerve stimulation in female larynx. A, Response of a type III fiber to single-stimulus pulses. No response is observed in response to a subthreshold stimulus. Suprathreshold stimuli result in an action potential. Calibration bars: 10 mV (vertical), 20 msec (horizontal). B, Response of a type III fiber to stimulus trains. An action potential is produced in response to each stimulus pulse within a train. The stimulus voltage was the same as that eliciting an action potential in A. Calibration bars: 10 mV (vertical), 20 msec (horizontal).
Fiber recruitment in males and females to nerve stimulation in a sex-typical manner. Three physiologically distinct fiber types have been described. Fiber types I and II are found predominantly in males, and type III is found predominantly in females. Both male-typical fiber types generate subthreshold potentials in response to nerve stimulation. The female-typical fiber type responds to nerve stimulation in an all-or-none fashion; subthreshold potentials are never observed.

**Fiber recruitment in males and females**

One goal of this study was to account, at the cellular level, for sex differences in muscle fiber recruitment (Tobias and Kelley, 1987). In males, laryngeal muscle fibers are recruited throughout a stimulus train, while in females, muscle fibers are recruited only at the beginning of a stimulus train. The result is dramatic potentiation of compound action potentials in males but not in females. The predominance of type III fibers in females might contribute to the constancy of compound action potential amplitude. Type III fibers respond in an all-or-none fashion to each pulse presented to the laryngeal nerve. Once threshold for generation of an action potential is attained, spikes are faithfully produced throughout a stimulus train. The coupling observed between female fibers might contribute to this result, since all coupled fibers would produce an action potential nearly synchronously. Male laryngeal muscle contains predominantly 2 fiber types: I and II. Type I fibers, like type III fibers, produce action potentials in response to each suprathreshold pulse within a stimulus train. Thus type I fibers contribute to the baseline amplitude of the compound action potential in males. In contrast to females, however, males also possess a large proportion of type II fibers, which are recruited as the stimulus train proceeds. Since type II fibers produce action potentials only in response to repetitive nerve stimulation, they add to the response generated by type I fibers, effectively increasing the compound action potential amplitude as the train proceeds. Each type II fiber produces an action potential after a different number of subthreshold events (see Fig. 2 for 2 examples). The frequency dependence of many type II fibers may also explain why compound action potential amplitude increases with stimulus frequency (Tobias and Kelley, 1987).

**Pre- versus postsynaptic contribution to muscle fiber type**

Both male laryngeal muscle fiber types (I and II) produce subthreshold potentials in response to nerve stimulation. Analysis of miniature endplate potentials in frog sartorius and cutaneous pectoris muscles reveals that individual synapses frequently produce only PSPs, not action potentials. Fewer quanta of neurotransmitter are released at low than at high safety factor junctions, consistent with differences in calcium availability within the nerve terminal (Grinnell and Herrera, 1980). Thus, the presence of subthreshold muscle responses can reflect a presynaptic property: release of small amounts of transmitter. In Xenopus laevis larynx, type II muscle fibers require repetitive stimulation by the laryngeal nerve for the production of action potentials. The increased size of PSPs during a stimulus train is probably due to facilitation, rather than to temporal summation, at the muscle fiber membrane. The latter mechanism could operate only if the interstimulus interval were less than the duration of a PSP, resulting in summation. In our experiments, the membrane potential always returned to resting level prior to the next response. Thus, we suggest that type II muscle fibers produce action potentials in response to increased transmitter release at the presynaptic terminal during repetitive nerve stimulation. Only one electrophysiologically identifiable fiber type has been described in female laryngeal muscle (type III). Type III fibers are typical of most vertebrate skeletal muscle: enough transmitter is released in response to each motor neuron action potential to guarantee a postsynaptic action potential. Apparent differences in transmitter release at the presynaptic terminal in males and females suggest that laryngeal motor neurons may also be physiologically different in the 2 sexes.

We have considered whether the detection of subthreshold events in male-typical fiber types only might result from sex differences in the location of neuromuscular junctions within the muscle, or from sex differences in muscle membrane properties. Karnovsky staining of laryngeal muscle has revealed that endplates are located in identical positions for males and females, the middle of the laryngeal muscle at all anterior/posterior levels (unpublished observations). This staining pattern is identical in both sexes. Since the recording electrode was randomly placed in the muscle in all experiments, electrode position probably does not account for sex differences in fiber type observed in this study. However, even if the electrode were the same distance from the endplate in both sexes, subthreshold events might be recorded only in males if the space constant of the membrane in males was larger. Fiber diameter and space constant are proportional; passive electrical events decay over a shorter distance in smaller fibers. Given the small diameter of most male laryngeal muscle fibers (Sassoon et al., 1987), it is unlikely that the smallest laryngeal muscle fibers are routinely, let alone exclusively, encountered in the female (see below).

**Lack of correlation between histochemical and electrophysiological fiber types**

Male laryngeal muscle is comprised of a homogeneous population of fast-twitch, highly oxidative (fatigue-resistant) fibers (Sassoon et al., 1987). In contrast, female laryngeal muscle is comprised of 3 histochemically identifiable fiber types: small, slow-twitch, highly oxidative (74%); medium-sized, fast-twitch, highly oxidative (6%); and large, fast-twitch, moderately oxidative (20%; Sassoon et al., 1987). We have described at least 2 electrophysiologically distinct fiber types in male larynx and one in female larynx. Thus, male larynx is homogeneous hist-

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**Figure 4.** Photomicrographs of Lucifer yellow-labeled fibers in laryngeal muscle. The entire larynx was transversely sectioned (25 μm). Labeled fibers could be reliably followed from section to section for distances up to 1 mm. Calibration bar, 30 μm. A: A single fiber is labeled following dye injection into an individual fiber in a male larynx. B: Multiple labeling observed following dye injection into an individual fiber in a female larynx. The fiber injected was within the superficial (i.e., lateral) third of the muscle leaflet shown in the photomicrograph. Muscle fibers of different diameters are dye-coupled.
androgen secretion. The presence of male-typical fibers in de-
elimination. Type I fibers would persist in males because of
innervated high safety factor synapse, then type I fibers in the
postmetamorphic development, males are exposed to rapidly
developing males at times when other skeletal muscles have
rodents (Jordan et al., 1988). In this muscle, multiple inner-
Testosterone-regulated synapse elimination has been postulated
development (Bennett and Pettigrew, 1975; Brown et al., 1976).
Angaut-Petit and Mallart (1979). The persis-
tically associated with multiply innervated endplates. The
absence of subthreshold potentials in type III fibers
does not prove that these fibers are singly innervated. In Xenopus
neurons. The absence of subthreshold potentials in type III fibers
response to androgen treatment (Sassoon and Kelley, 1986).
If synapse elimination is androgen-sensitive
in larynx, and if loss of multiple innervation results in a singly
eliminated supernumerary synapses. The muscle is lost post-
form. Since gap junctions are a common feature of embryonic
tissue, and have not been reported in adult skeletal muscle, the
formation may result in a larger insertion area onto the arytenoid
discs, producing more uniform and effective tension on the discs.
Dye-coupling in female laryngeal muscle may be due to gap
junctions between adjacent fibers. The low molecular weight
dye Lucifer yellow readily passes through gap junctions and has
been used to indicate the presence of these membrane special-
izations in Xenopus laevis (Warner et al., 1984). Alternatively,
muscle fibers may be dye-coupled through cytoplasmic bridges,
the remnants of incomplete myocyte fusion during develop-
ment. According to one view, 15 nm bridges between cells are
formed by the incorporation of "fusiform vesicles" prior to
fusion (Kalderon and Gilula, 1979). It is conceivable that com-
plete fusion is blocked in developing female laryngeal muscle,
resulting in coupled fibers. Gap junction formation has also been
reported immediately preceding muscle fusion in the developing
hindlimb of rats (Rash and Staehelin, 1974). In either case, dye-
coupling indicates electrical continuity of fibers. The conse-
quence for females is the synchronous contraction of many fi-
bers.
Gap junctions have been observed in adult muscle in response
specific endocrine conditions. For example, shortly before
parturition, rat uterine muscle becomes extensively coupled (Sims
et al., 1982). Gap junctions can be induced in myometrial tissue
by exposure to estrogens (Garfield et al., 1980). In the present
study, dye-coupling was not observed in any of 8 testosterone-
treated adult females examined. The presence of coupling in
female laryngeal muscle is probably not due to circulating es-
trogen, since this tissue does not contain estrogen receptor
(U. O'Dougherty, D. A. Sassoon, and D. B. Kelley, unpublished
observations). Adult female larynx does, however, contain an-
dergon receptor (Segil et al., 1987). Thus, it is more likely that
exogenously administered testosterone interferes with the main-
tenance or synthesis of gap junctions. Alternatively, gap junc-
tions may be physically pulled apart in response to muscle fiber
hypertrophy or to an increase in extracellular matrix, both of
which have been observed in female laryngeal muscle in re-
sponse to androgen treatment (Sassoon and Kelley, 1986).
The larynx is sexually monomorphic and female-like early in
development. In response to continuously rising levels of an-
drogen after metamorphosis, the male larynx gradually matures
into its distinctly male-like adult form (Lambdin and Kelley,
1986; Sassoon and Kelley, 1986). In contrast, the female lar-
ynx—in the absence of androgens—retains its less developed
form. Since gap junctions are a common feature of embryonic
tissue, and have not been reported in adult skeletal muscle, the
presence of dye-coupling between female laryngeal muscle fibers
supports the hypothesis that the female larynx is indeed a less
mature form of the male larynx. The finding that muscle fiber
electrophysiology is also sexually dimorphic suggests that this
property, too, may be controlled by androgens during develop-
ment. The physiological differences we report here may rely
on motor neuron differences that appear to become nonmalle-
able once innervation is complete.

Multiple innervation of laryngeal fibers; androgen and synapse
elimination
Male-typical type I fibers exhibit a subthreshold event in re-
sponse to a low-voltage stimulus delivered to the laryngeal nerve
and an action potential in response to a higher-voltage stimulus,
suggesting that these fibers are innervated by at least 2 motor
fibers. The presence of male-typical fibers in developing males at times when other skeletal muscles have
eliminated supernumerary synapses. The muscle is lost post-
naturally in females (Venable, 1966). During postnatal days 7–34,
testosterone treatment increases the number of multiply innerv-
ated terminals, as well as increasing the number of terminals per fiber. In Xenopus laevis, the multiply innervated type I fibers
are frequent in males, but extremely rare in females. During
postmetamorphic development, males are exposed to rapidly
rising levels of circulating androgen; females are not (Lambdin
and Kelley, 1986). If synapse elimination is androgen-sensitive
in larynx, and if loss of multiple innervation results in a singly
innervated high safety factor synapse, then type I fibers in the
female frog may be converted to type III fibers by synapse
elimination. Type I fibers would persist in males because of
androgen secretion. The presence of male-typical fibers in de-
veloping female larynx has yet to be tested.

Coupling between female muscle fibers
Female laryngeal muscle fibers were dye-coupled in 8/20 cases.
Dye-coupling was not observed in male laryngeal muscle in any
of 15 experiments. Adult female larynx contains approximately
one-eighth the number of muscle fibers found in adult males
(4000 versus 32,000; Sassoon and Kelley, 1986). A possible
functional advantage of coupling may be to increase the number of
muscle fibers contracting in response to each motor neuron
action potential. This effect would, in turn, insure sufficient
tension on the arytenoid discs to produce a sound. Coupled
fibers are organized in sheets, each fiber (except the most medial
one) positioned lateral and dorsal to its neighbor. This arrange-
ment may result in a larger insertion area onto the arytenoid
discs, producing more uniform and effective tension on the discs.

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