Advancing adult age is associated with a progressive decrease in skeletal muscle mass, strength and quality known as sarcopenia (reviews in Roubenoff et al., 2000; Doherty, 2003; Deschenes, 2004). Sarcopenia affects healthy, physically active subjects and the rate of muscle loss in humans has been estimated to range 1-2% per year past the age of 50 (Hughes et al., 2002). Therefore, sarcopenia represents a powerful risk factor for frailty, loss of independence and physical disability in elderly since it is associated with decreased functional performance, greater risk of falls, increased motor disability. Moreover, the reduction in muscle mass can be associated with osteoporosis (Szulc et al., 2005), altered thermogenic capacity (Wilson and Morley, 2003), insulin resistance, type 2 diabetes mellitus, dyslipidemia, and hypertension (Karakelides and Sreekumaran Nair, 2005). Understanding the mechanisms leading to sarcopenia represents therefore an essential step to prevent and combat illness and disability at advanced age.

The mechanisms underlying age-related skeletal muscle wasting and weakness are manifold and still remain to be fully elucidated (review in Ryall, 2008; Thompson, 2009); they may include denervation and reinnervation of motor units (Ansved and Larsson, 1989), decline in anabolic hormone concentrations (Tenover, 1997), increased concentrations of inflammatory mediators (Bruunsgaard et al., 2001), decrement in microvascular function and exercise tolerance (Payne and Bearden, 2006), impairment of proteolytic and autophagic pathways (Combaret et al., 2009), loss of satellite cells (Verdijk et al., 2007), depletion of myonuclei (Dirks and Leeuwenburgh, 2005).

Despite the extensive literature on sarcopenia and, in particular, on the debated role of myonuclei loss in fibre atrophy (Dirks and Leeuwenburgh, 2005, Dupont-Versteegden, 2005; Gundersen and
Bruusgaard, 2008), at present no data on the morpho-functional features of skeletal muscle nuclei in sarcopenia are available.

In eukaryotic cells, nuclear ribonucleoprotein (RNP)-containing structures are part of the transcription and splicing machinery. At electron microscopy, these structures are recognized as perichromatin fibrils (PF), perichromatin granules (PG), and interchromatin granules (IG) (Fakan, 2004; Spector, 1996). PF are the morphological equivalent of hnRNA transcription and co-transcriptional splicing, while IG represent a storage site for snRNP and non-snRNP splicing factors, and PG are involved in the storage and the nucleus-to-cytoplasm transport of mRNA (recently reviewed in Biggiogera et al., 2007). All these components have specific intranuclear location: PF and PG are at the periphery of condensed chromatin, and IG in the so-called interchromatin space (for a review, see Puvion and Puvion-Dutilleul, 1996). This is a necessary prerequisite for the maturation of nuclear RNAs, so that whenever transcription and/or splicing are altered, the organization, composition, and intranuclear location of RNP-containing structures are also affected (Biggiogera et al., 2004, 2007).

In this view, in the present study, by combining morphometry and immunocytochemistry at light and electron microscopy, we examined the fine structure of myonuclei as well as the distribution and amount of factors involved in RNA processing in skeletal myofibres of adult and old rats. As study system, we chose the biceps brachii and the quadriceps femoris muscles, which mostly contain fast type II fibres and are therefore largely affected by sarcopenia (Larsson et al., 1978; Shorey et al., 1998; Lexell, 1995).

**Materials and Methods**

Two adult (9 months of age) and two old (28 months of age) male Wistar rats were used. All animals were bred under controlled environmental conditions with a 12 h light/dark cycle, and fed ad libitum with a standard commercial chow. The experimental protocols comply with the guidelines of the Italian Ministry of Health as well as with internationally recognized guidelines. The rats were deeply anaesthetised with pentobarbital (50 mg/Kg i.p.) and then perfused via the ascending aorta with a brief prewash of 0.09% NaCl solution followed by 300 mL of a fixative solution containing 4% paraformaldehyde in 0.1 M phosphate buffer, pH 7.4 at 4°C. Biceps brachii and quadriceps femoris muscles were quickly removed and placed in the same fixation solution for 2 h at 4°C.

**Light microscopy**

After fixation, muscle samples were dehydrated with ethanol and embedded in paraffin wax. Five-µm-thick muscle samples were sectioned both transversally and longitudinally and then submitted to immunohistochemical procedures for fibre typing: sections were incubated overnight at 4°C with a mouse monoclonal antibody recognizing the heavy chain of skeletal fast fibre myosin (clone MY-32, Sigma-Aldrich, Buchs, Switzerland), then revealed with an Alexa 488 conjugated antibody against mouse IgG (Molecular Probes, Invitrogen, Milan). The sections were finally counterstained for DNA with 0.1 µg/mL Hoechst 33258 to label the myonuclei and to detect the occurrence of apoptosis based on chromatin morphology. Micrographs were taken in an Olympus BX51 microscope equipped with a 100 W mercury lamp under the following conditions: 330- to 385-nm excitation filter (excf), 400-nm dichroic mirror (dm), and 420-nm barrier filter (bf), for Hoechst 33258; 450- to 480-nm excf, 500-nm dm, and 515 nm bf for Alexa 488. Images were recorded with an Olympus Camedia C-5050 digital camera and stored on a PC by the Olympus software for processing and printing. Morphometrical evaluations were performed by using the software Image J (NIH, USA). The cross-sectional area of transversally cut fibres was measured on 100 myofibres per sample at 40X magnification. Moreover, the percentage of fast and slow muscle fibres was calculated on a total of 300 fibres per muscle. To evaluate nuclear density in fast fibres, the area of longitudinally sectioned myofibres was measured up to a total of about 15,000 µm² per muscle (mean value±SE: 15,072.21±15.01 µm²) at 40X magnification, myonuclei were counted and the nuclear density was expressed as the number of myonuclei/100 µm² of myofibre area. Finally, the percentage of apoptotic nuclei was evaluated, using a x20 objective lens, on a total of 25 microscope fields (at least 2000 myonuclei per muscle sample were counted).
Transmission electron microscopy

After fixation, muscle samples were washed in Sörensen buffer and subsequently in phosphate buffer saline (PBS), kept in 0.5 M NH₄Cl in PBS for 45 min to block free aldehydes, dehydrated with ethanol and embedded in LR White resin polymerized under U.V. light. Ultrathin sections were collected on Formvar-carbon coated nickel grids and used for morphometrical and immuno cytological analyses.

Morphometrical evaluations were made on micrographs (x11,000) of twenty myonuclei from longitudinally sectioned fibres per muscle by using a computerized image analysis system (AnalyzeSIS Image processing, Soft Imaging System GmbH, Muenster, Germany). The following parameters were considered: area of nuclei, nucleoli and fibrillar centres (FCs); percentage of nuclear area occupied by condensed chromatin; percentage of total FC, dense fibrillar component (DFC) and granular component (GC) area per nucleolus; PG density (PG/µm² of nucleoplasm); nuclear pore frequency (NP/µm of perimeter).

To investigate the fine distribution of some RNA transcription and processing factors, muscle sections were treated with the following probes: mouse monoclonal antibodies directed against the active phosphorylated form of RNA polymerase II (Research Diagnostic Inc., Flanders, NJ) or against the (Sm)snRNP (small nuclear RNP) core protein (Abcam, Cambridge, MA); rabbit polyclonal antibodies were used against DNA/RNA hybrid molecules (Testillano et al., 1994) specifically occurring in the transcriptional sites (Malatesta et al., 2008, 2009), and the cleavage stimulation factor CstF (Veraldi et al., 2001). Sections were incubated with the primary antibodies revealed by secondary gold-conjugated probes as described in Malatesta et al. (2009). As controls, some grids were incubated without the primary antibody and then processed as described above.

In situ hybridization for polyadenylated RNAs was carried out by using a biotinylated oligo d(T) (Sigma-Aldrich, Buchs, Switzerland) reacted with a gold-conjugated goat anti-biotin antibody (Aurion, Wageningen, The Netherlands) as reported in Cisterna et al. (2008). As controls, some grids were incubated in the absence of the probe, or pre-treated with 0.1% RNAase for 18 h.

To reduce chromatin contrast and selectively reveal nuclear RNP constituents, the sections were bleached by the EDTA method (Bernhard, 1969) and observed in a Philips Morgagni TEM operating at 80kV and equipped with a Megaview II camera for digital image acquisition.

Quantitative assessment of the immunolabelling was carried out by estimating the gold grain density over selected cellular compartments on sections treated in the same run. The surface area of nucleoplasm and nucleolus was measured on fifteen randomly selected electron micrographs (x22,000) from each animal by using a computerised image analysis system (AnalyzeSIS Image processing, Soft Imaging System GmbH). For background evaluation samples treated in the absence of primary antibody or pre-treated with RNAase were considered. Gold grains present over each selected compartment were counted and the labelling density was expressed as number of gold grains/µm².

Statistics

Results for each measured parameter were pooled according to the experimental groups and the means±standard error of the mean (SE) values calculated. Statistical comparisons were performed by the Mann Whitney U test (significance set at p≤0.05).

Results

Light microscopy

Morphometrical results are summarised in Figure 1. Fast myofibres (Figure 2 a-d) showed markedly smaller cross-sectional area in both biceps and quadriceps muscles of old rats compared to adult animals; slow fibres also were reduced in size in old rats, but the difference between the two groups was not statistically significant. The percentage of slow fibre was lower in both muscles, as expected. In old rats the percentage of slow fibre was higher than in adult animals, but the difference was not significant. Nuclear density in the fast fibres (Figure 3 a,b,f) was significantly higher in old than in adult rats, but the difference was not significant. Nuclear density in the fast fibres (Figure 3 a,b,f) was significantly higher in old than in adult rats, but the difference was not significant. Nuclear density in the fast fibres (Figure 3 a,b,f) was significantly higher in old than in adult rats, but the difference was not significant. 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To reduce chromatin contrast and selectively reveal nuclear RNP constituents, the sections were bleached by the EDTA method (Bernhard, 1969) and observed in a Philips Morgagni TEM operating at 80kV and equipped with a Megaview II camera for digital image acquisition.
Transmission electron microscopy

In all muscle samples myofibres contained multiple, elongated cell nuclei typically placed close to the sarcolemma. Myonuclei generally showed finely irregular border, condensed chromatin clumps at both the nuclear and nucleolar periphery, one roundish nucleolus characterized by a few FC, and abundant DFC and GC (Figure 4). In the nucleoplasm, all the usual RNP structural constituents involved in pre-mRNA transcription and processing - i.e., PF, PG and IG - were obvious. Morphological evidence of nuclear apoptosis was never observed in all muscle samples examined.

In both biceps and quadriceps muscles of old rats, myonuclei had smaller area and were characterized by a higher percentage of condensed chromatin, a higher density of PG and nucleoli with

Figure 1. Light microscopy morphometric parameters considered in biceps and quadriceps muscle myofibres of adult and old rats (means±SE). Asterisks indicate significantly different values.

Figure 2. Immunolabelled fast type II fibres in the biceps (a, b) and quadriceps (c, d) of adult (a, c) and old (b, d) rats. Slow (S) type I fibres are scarce. Note the smaller size of fast type II fibres in old animals. Bar: 30 µm.

Figure 3. Quadriceps muscle from an adult (a) and an old (b) rat; DNA was stained with Hoechst 33258. Bar: 50 µm. Very few apoptotic nuclei with typical chromatin condensation and nuclear fragmentation (arrows) were observed in both adult (c,d) and old (e) animals. Bar: 20 µm. The nuclear density is higher in both biceps and quadriceps muscles of old animals (f); asterisks show significantly different values.
smaller FC compared to adult animals (Figure 5). Conversely, there was no difference in either the nucleolar area or the percentage of DFC and GC in adult and old rats.

Immunocytochemical results showed similar distribution of polymerase II, DNA/RNA hybrid molecules, snRNPs, cleavage factor CstF and polyadenylated RNA in adult and old rats. In both animal groups, polymerase II was exclusively associated with PF (Figure 6 a,b); DNA/RNA hybrid molecules specifically labelled PF (Figure 6 a,b) and the nucleolar DFC (not shown); snRNPs were restricted to PF and, at a lesser degree, to IGs (Figure 6 c,d); CstF and polyadenylated RNA were located essentially on the PF and also on RNP tails rising from PG (Figure 6 e-h).

Quantitative evaluation of the immunolabelling (Figure 7) revealed lower nucleoplasmic density of polymerase II, DNA/RNA hybrid molecules and snRNPs in both biceps and quadriceps muscles of old rats; conversely, CstF and polyadenylated RNA were more abundant in the nucleoplasm of old than adult rats. No difference was found between the two age groups for nucleolar labelling obtained with any probes.

Figure 4. Myonuclei of quadriceps from adult (a) and old (b) rats. In old animals, condensed chromatin (ch) is more abundant and perichromatin granules are more numerous (inset) than in adult ones, whereas nucleoli (nu) show a similar aspect. Bars: 500 nm; inset 250 nm.

Figure 5. Electron microscopy morphometric parameters considered in myonuclei of biceps and quadriceps muscles of adult and old rats (means±SE). Asterisks show significantly different values.
Figure 6. Myonuclei from adult (a,c,e,g) and old (b,d,f,h) rats. a,b. Quadriceps muscles, anti-polymerase II (12 nm) and anti-DNA/RNA hybrid (5 nm) antibodies: both antibody probes specifically label perichromatin fibrils. c,d. Biceps muscles, anti-snRNP antibody: perichromatin fibrils and interchromatin granules (ig) are labelled. e,f. Biceps muscles, anti-CstF antibody: gold grains specifically label perichromatin fibrils. g,h. Quadriceps muscles, polyadenylated RNA: the labelling occurs on perichromatin fibrils and on RNP tails rising from perichromatin granules (arrows). Bars: 250 nm.
Background values were negligible in all immunolabelling experiments (*not shown*).

**Discussion**

Sarcopenia entails reduction of skeletal muscle mass, strength and resistance, with consequent impairment of performance in older individuals. In sarcopenia, muscles undergo loss of myofibres as well as decrease in total cross-sectional area of the remaining myofibres, both phenomena affecting preferentially type II fibres (Larsson et al., 1978; Shorey et al., 1998; Lexell, 1995). Accordingly, our data demonstrate a significant reduction in type II myofibre size in both biceps brachii and quadriceps femoris muscles of old rats. In addition, an increased - although not statistically significant - percentage of type I fibres is found in old animals, suggesting a preferential loss of type II fibres; however, there is conflicting evidence of changes in fibre type proportion in muscle in sarcopenia, some authors reporting no modifications (review in Hepple, 2003).

Muscle fibres are multinucleated syncitia and a reduction in fibre size could provoke alteration in nuclear number, nuclear domain size or both; however, studies on fibre atrophy conflict on whether these features undergo modifications or not. This is mainly due to the differences in muscle, myofibre and atrophy type that were investigated (for a recent review see Brooks et al., 2009); moreover, it has been demonstrated that myofibre size and nuclear number can be uncoupled in different phases of the ageing process (Wada et al., 2003; Bruusgaard et al., 2006). In the present study, it was found that nuclear density is higher in aged type II myofibres, according to previous reports on muscle atrophy (Tomonaga, 1977; Manta et al., 1987; Gallegly et al., 2004; Kadi et al., 2004; Edström and Ulfhake, 2005). This finding, together with the scarce presence of morphologically recognizable apoptotic nuclei (both at light and electron microscopy) suggests that only a limited nuclear death rate occurs in ageing skeletal muscles. It has been hypothesized that apoptotic induced loss of myonuclei may be involved in muscle protein degradation, thus contributing to fibre atrophy and, finally, myocyte death (Leeuwenburgh, 2003; Dirks and Leeuwenburgh, 2005; Alway and Siu, 2008). However, the role of apoptosis in myofibre depletion is still debated.

**Figure 7.** Quantitative immunoelectron microscopy of myonuclei of biceps and quadriceps muscles from adult and old rats; labelling density (gold grains/µm²) of several RNA processing factors in the nucleoplasm and nucleolar compartments (means±SE). Asterisks show significantly different values.
(Dirks and Leeuwenburgh, 2005, Dupont-Versteegden, 2005), some authors asserting that apoptotic signals mostly originate from satellite cells and connective tissue rather than from myofibres (Gundersen and Bruusgaard, 2008). The present observations are in agreement with previous data on hindlimb muscles of old mice demonstrating that apoptosis is quite scarce in both sedentary and trained animals (Zancanaro et al., 2007) as well as with ultrastructural studies showing the absence of typical apoptosis in atrophic myofibres (Rodrigues and Schmalbruch, 1995; Borisov and Carlson, 2000). However, it should be considered that our data have been collected in old animals where the sarcopenic process had already come to an advanced stage; this implies that a role of apoptosis in earlier phases cannot be excluded.

The myonuclei of aged type II fibres show morpho-functional features suggestive of a reduced activity in comparison to myonuclei of adult individuals. Smaller size and an increased amount of condensed chromatin together with lower quantity of phosphorylated polymerase II and DNA/RNA hybrid molecules, specifically occurring at transcriptional sites, clearly indicate a decrease in pre-mRNA transcription rate (Malatesta et al., 2008, 2009). The biological activity of many transcription factors is modulated by the redox state of the cell (Rojikind et al., 2002) and it is possible that the progressive increase of oxidative stress during ageing (Stadtman, 2004, 2006; Lesnefsky and Hoppel, 2006) contributes to the reduction of the transcription rate. According to the decrease in transcription rate, myonuclei of old rats contain lower amounts of nucleoplasmic splicing factors, involved in the co-transcriptional pre-mRNA splicing (Lührmann et al., 1990). In addition, myonuclei of aged fibres show an accumulation of cleavage factors and polyadenylated RNA, strongly suggesting a reduction in the processing and transport rate of pre-mRNA. The accumulation of PG further supports the hypothesis of decreased pre-mRNA transcriptional/co-transcriptional activities and reduced cytoplasmic export. In fact, PG represent storage and/or transport sites for spliced mRNA, and PG accumulation has been described as a consequence of altered pre-mRNA processing as well as of impaired intranuclear or nucleus-to-cytoplasm transport (e.g. Puvion et al., 1977; Puvion-Dutilleul et al., 1981; Lafarga et al., 1993; Zancanaro et al., 1993). Accordingly, a decrease in nucleus-to-cytoplasm transport factors (Pujol et al., 2002) has been reported during ageing, when the degradation systems also undergo alterations (Jameson, 2004) leading to the accumulation of crosslinked, insoluble and often oxidized proteins which may damage the intracellular transport mechanisms (Hallen, 2002; Kim et al., 2001); the intranuclear accumulation of RNA with poly(A) tails, known to be involved in stabilization, localization and translation of mRNAs (e.g. Soreq et al., 1974; Marbaix et al., 1975), could be also related to the reduced efficacy of the degradation machinery. Interestingly, malfunctions of the proteolytic and autophagic pathways are thought to be involved in sarcopenia (Attaix et al., 2005; Combaret et al., 2009).

On the other hand, myonuclear nucleoli in aged fibres do not show structural modifications suggestive of a marked activity decline. In fact, a significant reduction was observed in the FC only, whereas minor changes were found for the DFC and GC. However, the slight decrease of the DFC together with the lower (though not statistically significant) amount of DNA/RNA hybrid molecules in the nucleoli in aged fibres points to a reduction in the rRNA synthetic rate (Schwarzacher and Wachtler, 1993; Shaw and Jordan, 1995); in addition, the small increase of the GC may be the consequence of an impaired nucleus-to-cytoplasm transport of pre-ribosomal subunits.

It seems therefore that, during ageing, in myonuclei the entire production chain of RNA, from synthesis to cytoplasmic export, is less efficient. The reduction in muscle tissue mass with ageing has been attributed to a disruption in the regulation of skeletal muscle protein turnover, leading to a structural imbalance between muscle protein synthesis and degradation (reviewed in Koopman, 2009); in this view, altered RNA pathways may contribute to the reduced nuclear responsiveness to cellular metabolic needs. Accordingly, no decrease in the basal rates of muscle protein synthesis has been found in elderly, whereas during ageing the muscle protein synthetic response to the main anabolic stimuli, i.e. food intake and/or physical activity, is blunted (Koopman, 2009). Moreover, a reduction of nuclear activity would be consistent with the reduction in nuclear domain...
size observed in aged fibres. Interestingly, most of the changes found in myonuclei of aged fibres are reminiscent of those observed in senescent hepatocyte nuclei (Malatesta et al., 2003, 2004), and indicate that impairment in RNA pathways represents a general phenomenon during ageing. Moreover, defective RNA pathways due to nuclear sequestration of pre-mRNAs and processing factors have been recently related to diseases leading to muscle atrophy/dystrophy (Cardani et al., 2006; Wheeler and Thornton, 2007): the results of the present study suggest intriguing similarities in the basic mechanisms of sarcopenic and dystrophic processes, thus opening promising perspectives for preventing and treating skeletal muscle wasting.

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