Role of Ribonucleotides in Improving Muscle Cell Function

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Authors contributed equally to the study

Abstract

Based on data present in literature about some beneficial effects of nucleotides in reducing inflammation, and on lacking information about the mechanisms underlying the cellular aspects of exercise, the aim of this study was to examine the effects of the administration of a mixture of nucleotides and nucleotide-ribose on energy metabolism and contractility in cultured skeletal muscle cells in presence or absence of caffeine-dependent hyper-contraction. C2C12 cells were treated with various nucleotides forms, named Ribodiet®, RIBOCARE® MIX 1, RIBOCARE® MIX 2, and RNA-Na. Cell viability, calcium, and magnesium movements and the main kinases involved in glucose uptake have been analyzed. The level of glycogen and the presence of lactate have been evaluated as well. Since some important findings were obtained, additional experiments analyzed the effects of the same nucleotide forms under severe exercise conditions in order to understand the correct human use of nucleotides as food supplements to better support the exercise effects. The results obtained show for the first time, a direct effect of nucleotides on energy metabolism and calcium/magnesium movements during in vitro exercise simulation and provide useful indications to support muscle cells during exercise or during aging.

Keywords: Nucleotides; Muscle cells; Fatigue; Exercise physiology; Caffeine-induced contraction
1. Introduction

Fatigue is a feeling of extreme tiredness, which can result in a broad range of physical and mental unfitness [1, 2]. From the point of view of energy metabolism, it is characterized by the depletion of energy sources, the accumulation of end products, the decrease in glycemic levels, and glycogen consumption [3]. For this reason, efforts are needed to develop nutritional interventions capable of safely and effectively preventing adverse internal effects such as increased oxidative stress [4], and damage to mitochondria [5]. Interventions that reduce oxidative damage can effectively relieve fatigue, as suggested by the results of studies that have shown that the administration of antioxidant agents produced beneficial effects in mice [6, 7], also during the recovery phase after exercise [8]. In this context, the administration of nucleotides can represent a new intervention strategy, because they have demonstrated many beneficial functions, as both an ergogenic and immunostimulating additive including antitumor activity, immune modulation, protective capacity of the liver, normalization of metabolism, and improvement of exercise performance indices [9-13].

Additionally, nucleotides exhibit superior antioxidant and anti-aging properties, as demonstrated by a rat study [14].

Nucleotides are low-molecular-weight endogenously synthesized intracellular compounds, which form the basic units of the nucleic acids DNA and RNA, essential for cell division [15]. They are phosphoric nucleoside esters, made up of three components: a weakly basic nitrogenous compound, a pentose sugar, and one or more phosphate groups. The most important nucleotides are adenosine, guanosine, inosine, cytidine, and uridine monophosphates. Nucleotides are considered ‘semi-essential’ nutrients. This means that endogenous production meets the requirements in the basic maintenance condition, but exogenous administration is necessary for conditions such as childhood, stress, or tissue damage [16]. On the one hand, some tissues have a limited capacity for de novo nucleotide synthesis, and therefore require bases provided exogenously, on the other hand, some tissues regenerate more frequently than others and therefore require more nucleotides to support this activity [17]. For example, the intestinal mucosa, haematopoietic cells of the bone marrow, leucocytes, erythrocytes, and lymphocytes are not capable of de novo synthesis, and thus utilize the salvage pathway, suggesting that exogenous supply of nucleotides via the diet might be important for these cells [18]. On the other hand, individuals with a higher growth rate, such as pregnant women or children, also fall into the category of people with greater need, but these are not the only factors because also malnutrition or infections, injuries, damage to the tract intestinal (e.g. due to inflammatory bowel disease, IBD) may increase dietary needs. Indeed, nucleotide supplementation has also been linked to a reduction in IBD symptoms [19]. Moreover, nucleotides have a direct effect on the maintenance of intestinal mucosal integrity. It has been demonstrated that nucleotide supplementation in young rats increases the weight of the intestinal mucosa, villus height (by 25%), and the activity of brush border enzymes (maltase, sucrase, and lactase), suggesting an acceleration of gut growth and differentiation [20].

In addition, supplementation with a nucleoside-nucleotide mixture increases recovery after food deprivation, infection, or protein deficiency. Small intestine atrophy and decreased activity of brush border enzymes in rats quickly resolved with nucleotide supplementation [21]. As regards microbiota health status, in vivo studies show that dietary supplementation with nucleotides improves the intestinal flora and stimulates the growth of bifidobacteria [22]. Food nucleotides favor the development of the intestinal...
microbiota with a predominance of bifidobacteria and lactobacilli and a lower percentage of Gram-negative enterobacteria. Furthermore, nucleotides are mainly used in baby food as shown by many studies on healthy term infants where nucleotide supplementation is able to reduce the risk of diarrhea [18] or in a study about parenteral and baby milk formulas to improve intestinal health and to develop the immune system [20]. Moreover, another important field of application includes the immune system in which nucleotides have been shown to enhance capabilities, during cellular response to pathogens or viruses [23]. Nucleotides have also been shown to reduce the symptoms of common cold, including earache and sinus pain, and are especially effective when supplemented early in the infection [24].

Finally, as for immune system modulation, nucleotides influence both humoral and cell-mediated immunity: they accelerate T-cell-dependent antibody production and seem to exert actions on T-helper cells during antigen presentation, perhaps during cognitive cell-cell interactions. A nucleoside-nucleotide mixture stimulates the proliferation, differentiation, and maturation of neutrophils [20]. Nucleotides cause a transient increase in natural killer cell cytotoxicity, interleukin-2 production, and interferon-gamma secretion, and lower macrophage activation [25].

A recent application of nucleotide supplementation includes the effects on muscle cells in particular during fatigue and post-exercise recovery phase, although the intracellular mechanisms are poorly understood. Nucleotides are the building blocks for a variety of molecules, including ATP, which is broken down in the skeletal muscle cells to provide fuel for exercise. Nucleotides can be obtained from the diet and produced or retrieved by the body. Although nucleotides provide the building blocks of ATP, it has been suggested that sugar ribose limits the rate for synthesizing ATP. Previous studies have shown a decrease in ATP levels in the muscle after intense exercise, and this is linked to fatigue. Other studies have also reported benefits of nucleotide and/or ribose supplementation on exercise performance. However, no potential link between ATP metabolism and physical performance has yet been explored [26]. The main problem during exercise is to maintain adequate circulation and energy production but the main consequence includes free radical production that can lead to DNA damage or cell death. In this context the balance between oxidant and antioxidant levels is more important; if the antioxidant levels are low, the secondary effect can lead to inflammation and tissue damage. In this case, nucleotides are able to repair damaged DNA [27]. In addition, during the increase of muscle tone, nucleotides are essential for protein synthesis, and to reduce cortisol levels and increase secretory IgA [28].

The main important consequences of exercise and fatigue are skeletal muscle cramps that occur during or shortly following exercise in healthy individuals with no underlying metabolic, neurological, or endocrine pathology have been termed exercise-associated muscle cramps [29]. This condition can be present also in patients with several diseases influencing the quality of life [30]. Based on previous data on the use of ribose, the administration of nucleotide mixtures may be a possible new strategy to reduce or prevent exercise-induced muscle cramps and pain [31].

The aim of this study was to examine the effects of administering a mixture of nucleotides and nucleotide-ribose on fuel utilization and performance in skeletal muscle cells in the presence or absence of hyper-contraction caused by the presence of caffeine.
2. Material and Methods

2.1 Cell culture

Murine C2C12 myoblastic cell line was grown in Dulbecco’s Modified Eagle Medium (DMEM; Sigma-Aldrich, Milan, Italy) supplemented with 10% FBS (Sigma-Aldrich, Milan, Italy), 100 U/ml penicillin/streptomycin and maintained in an incubator at 37°C 5% CO₂. The cells were maintained at 40-70% density in order to have cell growth (cell proliferation) without cell differentiation. For experiments, cells were plated under different protocols: 1x10⁴ on 96-well plates to analyze the mitochondrial activity by MTT test; the 10 × 10⁴ cells/well in six-well plates in order to investigate the intracellular pathways activated by western blot and the Glucose/ Glycogen and lactate productions. The cell cycle stage of the subconfluence cells was synchronized by being cultured in DMEM for 24 h. The cells were then transferred to the growth medium at 2% FBS containing various nucleotides for a range time starting from 1h to 7h and 24h to verify the early phase of differentiation and the main mechanism activated. The nucleotides were tested in the presence or absence of high concentration of caffeine (2.5 mM) in order to mimic a condition leading to hypertrophy [32]. In particular, the nucleotides produced by Prosol Spa (Madone BG, Italy) tested in this study includes Ribodiet®, RIBOCARE® MIX 1, RIBOCARE® MIX 2, and RNA-Na.

2.2 Nucleotides mix

All nucleotides used to stimulate the cell were prepared by Prosol Spa (Madone BG, Italy). They are a pool of natural 5’-Nucleotides, obtained by enzymatic hydrolysis of yeast, named Ribodiet®, (5’-AMP.Na2, 5’-CMP.Na2, 5’-UMP.Na2, 5’-GMP.Na2, Nucleosides and other nucleotides); RIBOCARE® MIX 1 is a mix of five 5’ monophosphate nucleotides (5’ AMP free acid, 5’ CMP free acid, 5’ UMP free acid, 5’ GMP free acid and 5’ IMP free acid); RNA-Na is a ribonucleic acid, sodium salt, obtained from yeast by extraction without organic solvents and neutralized. All the mix were used without solvents and dissolved directly in the medium used to stimulate the cells at a final concentration of 100% (w/v) or 50% (w/v) which corresponded to human dosages of 0.5%/0.6% present in baby milk.

2.3 Cell viability

Cell viability was investigated after at the end of each stimulation time using MTT-based in vitro Toxicology Assay Kit (Sigma-Aldrich, Milan, Italy), as previously described [33]. Briefly, cells were incubated in DMEM without red phenol 0% FBS with 1% MTT dye for 2 h at 37°C in an incubator, and then the formazan crystals were solubilized by Solubilization Solution and the cell viability was determined by measuring the absorbance through a spectrometer (VICTORX4 multilabel plate reader) at 570 nm with correction at 690 nm. The results were obtained comparing them to control cells (100% viable).

2.4 [Mg2+]i Movements

Intracellular Magnesium [Mg2+]i was analyzed using Mg2+-sensitive fluorescent dye Mag-fura-2-AM (Furastra, Biotium) as previously described [33]. Briefly, the cells were incubated in a Hanks salt solution (Thermo Fisher Scientific, Waltham, USA) without Mg2+ but containing 10 mM glucose and supplemented with 20 mM HEPES/Tris (pH 7.4), 1.3 mM CaCl2, and 5 μM Mag-fura-2-AM at 37°C for 30 min. The fluorescence of cells loaded with Mag-fura-2 was monitored at regular intervals starting from 3min to 300min (15, 30, 60, 120, 150, 180, 210, 240, 270) with excitation at 340 nm and 380 nm. The acquisitions were obtained using a Fluorescence Spectrometer VICTOR X4 at the wavelength of emission of 510 nm with an
exposure time of 100 ms. The fluorescence ratios (340/380 nm) were calculated and compared to control; in Rmax and Rmin were analyzed by the addition of 50 mM MgCl2 and 100 mM EDTA, respectively [34].

2.5 [Ca2+] Movements
The analysis of calcium movements was performed following a classical technique [33]. Cells were washed twice with sterile PBS 1× and incubated with 5 μM Fura-2 AM (Sigma-Aldrich, Milan, Italy) for 30 min in the dark in PSS buffer without Ca2+ (1.5 mM KCl, 10 mM HEPES, 10 mM d-Glucose, 2 mM l-Glutamine, pH 7.4), in agitation at 37°C. After the stimulations with different ribonucleotides mix, the fluorescence was measured by the fluorescence spectrometer (VICTOR X4) at the wavelengths of 340 nm for excitation and 510 nm for emission. The results were expressed as a percentage compared to control cells.

2.6 Glucose uptake
After the presence of starved medium (DMEM without red phenol) overnight, cells were stimulated with Ribomix® in presence of 2-deoxyglucose (2-DG), followed by extensive washing three times with KrebseRinger’s buffer (20 mM HEPES, 5 mM KH2PO4, 1 mM MgSO4, 1 mM CaCl2, 136 mM NaCl, and 4.7 mM KCl, pH 7.4) following a classical protocol [35] and lysis in the Extraction buffer by several freeze/thaw cycles and then heat at 85°C for 40 min. The cooling cell lysate was put on ice for 5 minutes, neutralized by adding 10 μL of Neutralization Buffer, spun down at 13,000 × g to remove insoluble material and then the amount of glucose in the lysate was measured using Glucose Uptake Colorimetric Assay Kit (Sigma-Aldrich) was performed following the manufacturer’s instruction. After being reacted with mixed reagents A and B, the absorbance of the mixture was measured at 412 nm by spectrophotometer (VICTOR X4). Data were expressed as mean ± SD of pmol/μl derived by standard curve.

2.7 Glycogen measurement
The glycogen synthesis assay was performed as described in the literature [36] and following the manufacturer’s instruction (BioVision, Life Research, Scoresby Victoria, Australia). C2C12 cell lysates were collected and homogenates and boiled for 5 min to inactivate enzymes. Samples were then centrifuged at 13000 rpm for 5 min and the supernatant was pick up to induce the hydrolysis of glycogen to glucose in presence of an OxiRed probe to generate color (lmax = 570 nm). The glycogen concentration in the samples was calculated by C = Ay/Sv where Ay is the amount of glycogen (mg) in the sample as determined from a standard curve and Sv is the sample volume (ml) [36].

2.8 Lactate measurement
As reported in the literature [37] the quantification of lactate level on C2C12 cell was analyzed after lyses on supernatants using a lactate assay kit (BioVision) according to the manufacturer’s protocol. The contents of pyruvate and lactate were normalized to the protein amount.

2.9 ERK Activation Assay
ERK/MAPK and PI3K/Akt activities were measured by the InstantOne™ ELISA (Thermo Fisher, Milan, Italy) cell lysates following the manufacturer’s instructions [38]. Briefly, 50 μL/well of each lysed were tested in InstantOne ELISA microplate strips after 1 h at room temperature on a microplate shaker with the antibody cocktail. In the end, the detection reagent was added for 20 min and then stopped by adding a stop solution. The strips were measured by a spectrometer (VICTOR X4 multilabel plate reader) at 450 nm. The results were expressed as mean absorbance (%) compared to control.
2.10 Phospho-p38/MAPK ELISA test
After stimulations with all nucleotides in the presence or absence of 2.5 mM caffeine, cells were lysed to assess phosphorylation levels of p38 MAPK using a cold Lysis Buffer kit supplemented with protease and phosphatase inhibitors, following the manufacturer’s instruction (p38 MAPK alpha (pT180/pY182) + total p38 MAPK alpha ELISA Kit, Abcam, ab221012) [39]. Absorbance was read at 450 nm on a microplate reader and the results expressed as means (%) vs control.

2.11 Western blot analysis
At the end of each treatment, C2C12 cells were washed twice with ice-PBS 1x and lysed in ice Ripa Buffer (50 mM Hepes, 150 mM NaCl, 0,1% SDS, 1% TRITON 100x, 1% deoxycholate acid, 10% glycerol, 1.5 mM MgCl2, 1 mM EGTA, 1 mM NaF; all purchased from Sigma-Aldrich) supplemented with 2 mM sodium orthovanadate (Sigma-Aldrich), 1 mM phenylmethanesulfonyl fluoride (PMSF; Sigma-Aldrich) and 1:100 mix Protease Inhibitor Cocktail (Sigma-Aldrich). 40 μg proteins of each sample were resolved into 8% and 15% SDS-PAGE gels, and polyvinylidene difluoride (PVDF) membranes (GE Healthcare) were incubated overnight at 4°C with specific primary antibody: anti-Cyclin D1 (1:1000, Cell Signalling), anti-Desmin (1:1000, Santa-Cruz), anti-Phospho-AMPK (1:1000, Santa-Cruz), anti-AMPK (1:500, Santa-Cruz), and anti-SMA (1:1000, Santa-Cruz). Protein expression was normalized and verified through β-actin detection (1:5000; Sigma-Aldrich) and expressed as a mean ± SD (% vs control).

2.12 Statistical analysis
Data reported were obtained from at least four independent experiments running on triplicates for each experimental protocol and processed using Prism GraphPad statistical software. Results are expressed as mean ± SD using One-way ANOVA followed by Bonferroni post hoc test for statistical analysis. p values <0.05 were considered statistically significant.

3. Results
3.1 Time-dependent study of cell viability and proliferation on C2C12 Cells
In order to explore the intracellular mechanisms activated by nucleotides, firstly the analysis of mitochondrial health by MTT was carried out. The first set of experiments was performed to demonstrate the effect exerted on cell viability by direct exposure to different nucleotides preparations during the time (ranging from 1h to 7h) and to exclude any cytotoxic effects of the formulations. As shown in Table 1, all nucleotides were able to induce an increase on cell viability during the time; in particular Ribodiet®, Ribocare® MIX 1 and 2, RNA-Na-E showed a time-dependent increase on this parameter with a greater effect starting from 5h to 7h (p<0.05 vs control). Furthermore, the nucleotides RNA-Na-E, which was composed only of RNA, was able to induce a late response on cell viability, supporting the hypothesis of a late effect with respect to other nucleotides. All these data confirmed that none of the forms of nucleotides had cytotoxic effect overtime on muscle cells.

In addition, in order to understand if these nucleotides were also able to induce a physiological differentiation on myotubes of C2C12 cells, additional experiments were carried out stimulating cells for 24h analyzing their proliferation rate. As reported in Figure 1A, the effects of all nucleotides lead to a stable value around 24h of stimulation on cell viability (p<0.05 vs control), since the main effect was observed by Ribocare® MIX 2 stimulation compared to other nucleotides (about 5 round compared to Ribodiet® p<0.05; about 25% compared to Ribocare® MIX 1; about 40% compared to RNA-Na-E). This finding indicated for the first time that nucleotides were able to induce the proliferation
and differentiation phases, as confirmed by cyclin D1 and desmin on western blot analysis (Figure 1B and 1C). Indeed, as illustrated in Figure 1B, nucleotide Ribocare® MIX 1 was able to induce a greater effect compared to other nucleotides (p<0.05) confirming that the expression cyclin D1 is abundant in proliferating myoblasts. Similar data were reported by desmin analysis (Figure 1C) in which nucleotide Ribocare® MIX 1 showed the ability to induce greater expression compared to all other nucleotides (p<0.05), confirming a better influence on proliferation of this form.

3.2 Nucleotide-induced contraction mechanisms on C2C12 cells

As is known, muscle contraction during exercise depends on variations in the intracellular concentrations of calcium ions (Ca2+). Given the behavior of calcium in C2C12 cells is similar to that in vivo, changes in cytosolic calcium and magnesium levels in all treatments using Fura-2 and Furaptra, respectively, were studied in these cells. As shown in Figure 2, the administration of all nucleotides was able to modify the calcium movements in a few minutes and this rapid effect was followed by a reduction in intracellular magnesium, supporting the hypothesis of a physiological role exerted by nucleotides during muscle contraction. This intracellular ion flow cycle was maintained throughout the analyzed period (from 3 'to 300'), indicating a rapid phase of contraction and relaxation, as occurs in the human physiological mechanism.

In order to clarify the balance between the calcium and magnesium movements, a schematic table (Table 2) summarizes the main effects over time of all the nucleotides tested. In particular, stimulation with Ribodiet® was able to maintain a physiological trend compared to other treatments; Ribocare® mix 1 seemed to support exercise better than other nucleotides, inducing rapid contraction of muscle cells; Ribocare® mix 2 appeared to have a similar tendency to Ribocare® mix 1 but its effect was slower than Ribocare® mix 1 (at 30 min). And it confirmed their ability to have a slow effect on muscle cells that have a long relaxing phase. However, it is currently impossible to determine which nucleotide is capable of best supporting exercise.

3.3 Analysis of the intracellular pathways activated by nucleotides on C2C12 cells

To explore whether different nucleotides can support different phases and moments in the intracellular mechanisms involved in an exercise, further experiments were performed by analyzing some important kinases in muscle cells. In particular, the phosphorylation of PI3K/Akt, ERKs/MAPK and 5′ AMP-activated protein kinase (AMPK) which are intracellular signaling proteins typically activated in exercised/contracted skeletal muscle, were investigated. As reported in Figure 3, the activity assay of ERKs/MAPK was analyzed at different times (at 3min and 24h), in order to explore the rapid effects and upregulation caused by a long incubation period. Data reported confirmed its role to help the increases of glucose transport in few mins of stimulation induced by both Ribodiet® and Ribocare® MIX 1 nucleotides compared to Ribocare® MIX 2 and RNA-Na-E (p<0.05) and the greater effects was observed by Ribodiet® (p<0.05 vs all nucleotides), indicating a rapid activation of the intracellular pathway leading to glucose uptake supporting the early phase of exercise. The reported data confirmed the role of ERKs/MAP in facilitating the increase in glucose transport after a few minutes of stimulation induced by both Ribodiet® and Ribocare® MIX 1 nucleotides compared to Ribocare® MIX 2 and RNA-Na-E (p <0.05). The greatest effects were observed with Ribodiet® (p <0.05 vs all other nucleotides), indicating rapid activation of the intracellular pathway leading to rapid glucose uptake to
support the initial phase of exercise. In contrast, the 24-hour effects support previous data on the proliferation phase and all nucleotides were able to induce ERK activity. However, the Ribocare® MIX 2 showed better influence than other nucleotides (p <0.05). A second major pathway supporting GLUT4 translocation and glucose uptake, PI3K / Akt (Figure 3B) at 24 hours, confirmed its ability to support a late phase of cell activation, since only formulations Ribocare® MIX 1 and Ribocare® MIX 2 were able to induce its activation (p <0.05). Furthermore, Ribocare® MIX 2 nucleotide induced a more potent effect (approximately 5 times compared to Ribocare® MIX 1, p <0.05). Another key regulator of cellular metabolism which plays a predominant role in catabolic mechanisms: glucose transport and differentiation mechanism of myotubes is AMPK, as reported in Figure 3C at 24h. Nucleotides Ribodiet®, Ribocare® MIX 1 and RNA-Na-E were able to induce greater effects compared to Ribocare® MIX 2 (p<0.05).

Since the exercise in vivo induces a cascade of intracellular signaling (e.g. activation of AMPK), proteomic (e.g. increased GLUT4 expression) and metabolic changes (e.g. enhanced glucose uptake) within skeletal muscle cells additional parameters such as glucose uptake, glycogen consumption and lactate were consequently investigated. As reported in Table 3, the intracellular mechanism reflects what observed by glucose and glycogen analysis; indeed, a rapid glucose uptake, increases glycogen accumulation and increases the rate of extracellular lactate following a physiological mechanism of exercise. In particular, in presence of Ribodiet®, Ribocare® MIX 2 and RNA-Na-E the glucose uptake was significantly increased compared to control (p<0.05). Among these, the best effect seems to be exerted by Ribocare® MIX 2 compared to Ribodiet® (about 11.22%) and RNA-Na-E (about 2.5 times). Glycogen, which is the main intracellular store of glucose, was increased by the stimulation of the cells with all tested nucleotides (p<0.05), indicating that they increased the uptake of glucose to feed the glycogen synthesis process and the main effect was shown by nucleotide Ribocare® MIX 2 (p<0.05 compared to Ribocare® MIX 1 and RNA-Na-E). Finally, lactate, which has an important role in energy metabolism, measured in extracellular environment showed an increase after stimulation with all nucleotides compared to control (p<0.05) and Ribocare® MIX 2 revealed a main effect compared to other nucleotides indicating that this form may be the better choice to support the muscle exercise.

3.4 In vitro exercise simulation by caffeine-induced contraction

It is well established by in vitro studies that caffeine has a direct effect on muscle contraction. It acts on the sarcoplasmic reticulum, increasing calcium permeability and making it readily available for the contraction mechanism [40]. Caffeine treatment has been applied to C2C12 cells with the specific intent to provide an ‘exercise-like treatment’, mimic exercise activation to examine exercise-regulated changes in signal transduction and metabolism. Cells were pre-treated with 2.5 mM caffeine and then treated with all nucleotides in various time points. In order to understand the effects on contraction mechanisms, the levels of calcium and p38/MAPK activity were analyzed. As reported in Figure 4A, all nucleotides tend to dampen calcium oscillations overtime to reach the relaxation phase in the presence of caffeine; indeed, below 2h it was observed all the forms were in resting, at 30 min they tend to contraction again while at 1h the system is rebalanced again (relaxation). This effect was confirmed by the analysis of p38 activity by assay kit (Figure 4B), in which a reduction of contraction was observed starting from 3min up to 2h and then the activity returns to physiological levels starting from 3h.
up to 5h. In particular, the addition of nucleotide Ribodiet® counteracts caffeine-induced hypercontraction with a better effect compared to Ribocare® MIX 1 (p<0.05). These data demonstrate that nucleotides have different properties during muscular contraction; so that in severe contraction Ribocare® MIX 2 plays a minor role compared to Ribodiet®.

In order to confirm these effects, the analysis of glucose, glycogen and lactate was done after pretreatment with caffeine and then stimulated with all nucleotides. As shown in Table 4, caffeine alone induced severe contraction accompanied by high glucose uptake and glycogen and a significant reduction on extracellular lactate (p<0.05), indicating a negative condition of the cells which are in hypercontractility state. The addition of single nucleotides was able to reduce this negative condition, leading the extracellular lactate to the control value (p<0.05 vs caffeine alone). Finally, Ribodiet® confirmed to be able to support muscle activity in aerobic physiological conditions.

Since the severe contraction can cause muscle hypertrophy accompanied by cell death when the stress condition is maintaining for a long time, additional experiments were carried out at 24h analyzing cell viability, ERK/MAPK activity and SMA proteins expressions to verify if nucleotide may be also helpful to restore the damage activating cell proliferation. As reported in Figure 5A, a high dose of caffeine caused a decrease in cell viability (p<0.05 vs control) confirming data present in literature and this negative effect was counteracted by all tested nucleotides (p<0.05). However, both Ribodiet® and Ribocare® MIX 2 seem to have greater effects than the other forms, where the former was also able to reduce hyper-contractility and the latter did not counteract the movement of the calcium during contraction, but induced cell proliferation, which could activate survival mechanisms. These findings have been confirmed by ERK activity assay (Figure 5B), in which only Ribodiet® and Ribocare® MIX 2 were able to have the main effects compared to other nucleotides (p<0.05). Furthermore, all nucleotides were able to counteract the negative effect of caffeine alone (p<0.05). Finally, the analysis of regeneration and myogenesis markers such as smooth muscle actin (SMA) showed similar results (Figure 5C). indeed all nucleotides counteracted the negative effect of caffeine and the main effects were observed with Ribodiet® and Ribocare® MIX 2 stimulations (p<0.05) compared to other nucleotides while between Ribodiet® and Ribocare® MIX 2 no significant change was observed, indicating an equal beneficial action on muscle cells in severe contraction conditions. All of these results support the hypothesis that different nucleotides may have different roles during different phases of muscle activity and also reveal different beneficial effects during a strong contraction.

<table>
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<td>3.23 ± 3.02</td>
<td>1.33 ± 0.57</td>
<td>5.09 ± 1.01</td>
<td>5.1 ± 1.01*</td>
<td>10.04 ± 1*</td>
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<td>1 ± 0.9</td>
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<td>1.35 ± 0.8</td>
<td>55.04 ± 2*</td>
<td>30.24 ± 1.08*</td>
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Table 1: Cell viability of C2C12 cells stimulated with different nucleotides forms. The data reported are expressed as means ± SD (%) compared to control value (0%). * p<0.05 vs control.
(A) Cell viability measured by MTT test; (B) and (C) Cyclin D1 and desmin expressions measured by western blot, respectively. Data reported are expressed as means ± SD (%) compared to control (0 line) of 4 independent experiments. *p<0.05 vs control; the bars p<0.05 between different nucleotides.

**Figure 1:** Cell viability and proliferation analysis at 24h of stimulation.
(A) Calcium and (B) Magnesium flux measured by Fura and Furaptra fluorescence dyes respectively during time (starting from 3 min to 5h). Data reported are expressed as means ± SD (%) of four experiments vs control (0 line).

**Figure 2:** Analysis of calcium and magnesium flux on C2C12 cells.

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<td>C</td>
<td>S</td>
<td>R</td>
<td>R</td>
</tr>
</tbody>
</table>

C=contraction; R=relaxation; S=resting phase.

**Table 2:** Analysis of the balance between calcium and magnesium flux under nucleotides stimulations on C2C12 cells. The empirical analysis summarized the effects observed on Figure 1 and the conclusion of each time point was obtained by the ratio from calcium and magnesium (who is big the other correspond to the main effect observed).
Figure 3: ELISA activity and western blot analysis. (A) and (B) ELISA test of ERKs/MAPK activity measured at 3 min (on the left) and 24 h (on the right) and PI3K/Akt at 24 h measured after stimulation with all nucleotides tested. Data are expressed as means ± SD of four experiments compared to control (0% line); (C) Western blot and densitometric analysis of the AMPK expression. The images shown are an example of each protein from four independent experiments. Data are expressed as means ± SD (%) normalized on total protein and verified on β-actin detection.
<table>
<thead>
<tr>
<th></th>
<th>Glycogen microg/microl</th>
<th>Extracell. Lactate nmol/microl</th>
<th>Glucose pmol/microl</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTR</td>
<td>0.639 ± 0.01</td>
<td>142 ± 0.03</td>
<td>36.27 ± 1.62</td>
</tr>
<tr>
<td>Ribodiet®</td>
<td>0.745 ± 0.01*</td>
<td>142.1 ± 0.02*</td>
<td>42.42 ± 2.66*</td>
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<tr>
<td>Ribocare® Mix 1</td>
<td>0.692 ± 0.02*</td>
<td>141.2 ± 0.03*</td>
<td>38.09 ± 1.65</td>
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<tr>
<td>Ribocare® Mix 2</td>
<td>0.752 ± 0.03*</td>
<td>143.7 ± 0.04*</td>
<td>47.18 ± 3.02*</td>
</tr>
<tr>
<td>RNA-Na-E</td>
<td>0.693 ± 0.01*</td>
<td>141 ± 0.02*</td>
<td>18.30 ± 2.07*</td>
</tr>
</tbody>
</table>

**Table 3:** Analysis of Glucose uptake, glycogen and lactate by ELISA test. Data reported are expressed as means ± SD of four independent experiments carried out in triplicates. CTR= control sample. *p<0.05 vs control.

<table>
<thead>
<tr>
<th></th>
<th>Glycogen microg/microl</th>
<th>Extracell. Lactate nmol/microl</th>
<th>Glucose pmol/microl</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTR</td>
<td>0.66 ± 0.024</td>
<td>143.6 ± 1.51</td>
<td>34.27 ± 2.05</td>
</tr>
<tr>
<td>Caffeine + Ribodiet®</td>
<td>0.93 ± 0.01*</td>
<td>144.3 ± 0.06</td>
<td>30.97 ± 0.12*</td>
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<tr>
<td>Caffeine + Ribocare® Mix 1</td>
<td>0.64 ± 0.5</td>
<td>145.3 ± 0.62</td>
<td>30.50 ± 0.3</td>
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<tr>
<td>Caffeine + Ribocare® Mix 2</td>
<td>0.129 ± 0.11*</td>
<td>146.9 ± 1.78</td>
<td>20.54 ± 1.33*</td>
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<tr>
<td>Caffeine + RNA-Na-E</td>
<td>0.74 ± 0.5*</td>
<td>143.1 ± 0.1</td>
<td>25.53 ± 0.16*</td>
</tr>
<tr>
<td>Caffeine</td>
<td>2.08 ± 0.03*</td>
<td>120.7 ± 2.53*</td>
<td>66.73 ± 2.61*</td>
</tr>
</tbody>
</table>

**Table 4:** Analysis of Glucose uptake, glycogen and lactate by ELISA test. Data reported are expressed as means ± SD of four independent experiments carried out in triplicates. CTR= control sample. * p<0.05 vs control.
(A) Calcium fluorescence and (B) p38 activity. Data are expressed as means ± SD (%) compared to control value (0 line) of four independent experiments.

**Figure 4:** Calcium movement and p38/MAPK activity analyzed in presence of 2.5 mM caffeine.
Figure 5: Cell viability and protein activated by nucleotides in presence of caffeine. (A) Cell viability measured by MTT test; (B) ERK/MAPK activity measured by ELISA test; (C) Western blot and densitometric analysis. Data reported are expressed as means ± SD of four independent experiments expressed as % vs control (0 line). Western blot reported is an example and Data are expressed as means ± SD (%) normalized on total protein and verified on β-actin detection. * p<0.05 vs control; ** p<0.05 vs caffeine; the bars p<0.05 between different nucleotides.
4. Discussion
In this study, conducted on muscle cells, the intracellular mechanisms evoked by the main nucleotides used as food supplements were verified for the first time. The data obtained made it possible to identify for the first time two important roles they played; in physiological conditions, the tested nucleotides can support and improve the energy balance of the cell and in conditions of intense physical exercise they support a prevention mechanism capable of countering the damage caused by severe hypercontraction conditions. In particular, different effects of Ribodiet®, Ribocare® mix 1 and 2 were observed with respect to RNA-Na. In fact, in physiological operating conditions starting from the analyzed parameters Ribocare® MIX 2 was found to have a greater effect than other nucleotides. However, Ribodiet® seems to play a major role during the first phase of the exercise compared to Ribocare® MIX 2. On the other hand, Ribocare® MIX 2 and Ribodiet® play a similar role under intense exercise conditions by activating different damage restoration mechanisms (both SMA protein and cell differentiation).

A strength of this study was the use of an in vitro model of physical exercise [41], through the administration of caffeine, which allowed to investigating the cellular effects of individual aspects of the physiology of exercises, such as the contraction of the skeletal muscle or the activation of signaling pathways that respond to exercise, directly on skeletal muscle cells, as well as exercise-mediated cross-talk between different cell types [42]. C2C12 cells have been extensively used to study the effects of various agents not only in myoblasts, which are an undifferentiated state of myogenic differentiation, but also in myotubes, a differentiated state on insulin-dependent or independent mechanisms of glucose uptake and expression of GLUT-4 [43].

Skeletal muscle is an absolutely important element for relational life as it supports physical activity that allows us to interact with the environment around us [44] and it is the largest tissue in the human body, since it represents up to 40-50% of body weight [43]. Although there has been a lot of in-depth research on exercise in recent years, the many human health benefits of controlling muscle tissue still remain to be fully understood. Such benefits could be improved glucose disposal, changes in muscle fiber types, induction of angiogenesis, and regulation of the immune system in both the prevention and treatment of various chronic diseases [45, 46]. However, over the course of a lifetime, the trophism of muscle tissue can undergo profound changes, such as skeletal muscle hypertrophies that occur following intense resistance training or atrophies that occur rapidly, for example during prolonged bed rest [47-49]. Also, another important factor is that skeletal muscle mass and strength decrease with age, which can lead to decreased levels of physical activity and increased risk of falling with consequent negative effects that affect the quality of life [44]. Moreover, the exact intracellular mechanisms responsible for the benefits of exercise remain to be determined. Indeed, skeletal muscle contraction per se likely contributes to some or many of the exercise benefits, be that through direct effects within the contracting skeletal muscle cells, such as improved glucose uptake and insulin sensitivity [50, 51] or mediated by factors released from the skeletal muscle cells, such as myokines and/or metabolites [52, 53]. Determining key mechanisms and signal transduction specific to each exercise-triggered benefit could optimize its beneficial role in maintaining health, minimizing impairment of exercise adaptation, and could facilitate the identification of novel drug targets responsive to exercise. In this context, many important data on the role of nutrition in the regulation of skeletal muscle mass are reported in the literature. Recent
studies have reported the importance of nutrition-induced regulation of mRNA levels for a range of metabolic genes [44]. For example, Anthony et al. reported that the oral administration of leucine stimulated protein synthesis in skeletal muscles [54], or Yoshihara et al. that the administration of 6% beef extract for 4 weeks increased leg muscle weight and modified the muscle fiber type [55]. Furthermore, the consumption of dietary supplements with “ergogenic” claims occurs in numerous target categories, such as athletes [56], the elderly [57], patients with chronic diseases [58], and sedentary adults [59]. The influence of plants-based food supplements in improving exercise performance has been a focus of many studies, in order to explain the main mechanism of every single ingredient. Indeed, Antioxidant phytochemical supplements are used by physically active individuals as ergogenic functional foods for improving physiological or metabolic responses that may increase performance during exercise [60]. High-intensity exercise training leads to excessive levels of reactive oxygen species, pro-inflammatory cytokines, and excess concentration of cortisol which may hamper sports performance, induce fatigue, and delay recovery [61]. Therefore, developing antioxidant phytochemicals to prevent exercise-induced oxidative stress along with improving adaptation responses to endurance training is an attractive strategy for physically active individuals [62]. Thus, it is necessary to study in-depth the ever-growing area of nutrients/nutraceuticals regarding various aspects of skeletal muscle health and associated cellular mechanisms to formulate evidence-based recommendations [63]. In this context, our study may suggest a possible new intervention strategy through dietary supplementation capable of acting on physical exercise, using a natural compound, nucleotides. The use of nucleotides is characterized by a greater compliance for muscle cells, since they are very important for mitochondrial function by promoting many functions of muscle cells. C2C12 cells, the mouse skeletal muscle cell line, were used for this. Although in vivo experiments with human volunteers represent a more integrated model for examining responses to exercise, a preemptive examination of the mechanisms from a molecular point of view is needed to optimize experimental models to resemble real elements of the physiology of exercise, but with a more reductionist approach [42].

5. Conclusion
In conclusion, this study lays the foundations for the development of a new strategy, based on nucleotides dietary supplements, which can provide beneficial effects on muscle function, whether during physical exercise or when recovering from an exercise or to counteract the physiological process of muscle mass loss in the elderly.

Statement of Competing Interests
The authors have no competing interests.

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References


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