In muscular dystrophies, muscle membrane fragility results in a tissue-specific increase of danger-associated molecular pattern molecules (DAMPs) and infiltration of inflammatory cells. The DAMP extracellular ATP (eATP) released by dying myofibers steadily activates muscle and immune purinergic receptors exerting dual negative effects: a direct damage linked to altered intracellular calcium homeostasis in muscle cells and an indirect toxicity through the triggering of the immune response and inhibition of regulatory T cells. Accordingly, pharmacologic and genetic inhibition of eATP signaling may provide a therapeutic approach to slow disease progression in α-sarcoglycanopathy. (Am J Pathol 2019, 189: 354–369; https://doi.org/10.1016/j.ajpath.2018.10.008)
the first decade of life and rapid evolution, to milder forms with later onset and slower progression. Currently, no treatment is available for these patients.

Sarcoglycans and dystrophin are main components of the dystrophin–glycoprotein complex, the crucial structural linkage between the intracellular cytoskeleton and the extracellular matrix that provides stability to the sarcolemma and offers muscle fibers protection from muscle contraction–induced stress. In both sarcoglycanopathies and dystrophinopathies, dystrophin–glycoprotein complex disruption leads to a mechanically weaker muscle plasma membrane that is easily damaged during contraction, allowing release of intracellular molecules, which act as damage-associated molecular pattern molecules (DAMPs). These self-molecules execute precise intracellular tasks in both innate and adaptive immunity, exerting disparate functions when released into the extracellular space. Among DAMPs, extracellular ATP (eATP) and its derivatives are pleiotropic regulators of cell function. In muscle, the primary function of ATP is energy transfer; however, after cell damage, it is quickly released in the extracellular space, where it acts as DAMP. Once in the extracellular space, eATP activates plasma-membrane nonselective ion channels purinergic P2X1 to 7 receptors and P2Y1, 2, 4, 6, and 11 to 14, guanine nucleotide binding protein-coupled receptors. The role of eATP and P2X purinergic signal on mdx dystrophopathy has been recently well established. We have shown that blockade of eATP/P2X purinergic signaling by periodate-oxidase ATP (oATP) delayed the progression of the dystrophic phenotype, dampening the local inflammatory response in dystrophin-deficient mdx mice. These results are in agreement with the phenotype displayed by mdx/P2X7 double-knockout mice, which showed a reduction of the inflammatory and fibrotic response in skeletal muscles and heart.

We hypothesized that the role of eATP may be equally relevant in α-SG deficiency because this protein displays in its extracellular domain an ecto-ATPase activity, which allows conversion of ATP to the inactive form ADP. The role of eATP and P2X purinergic signal on α-SG deficiency in LGMD2D patients.

Materials and Methods

In Vivo Treatment with oATP

Sgca knockout mice [kindly donated by Prof. Giulio Cossu (University of Manchester, Manchester UK)] were treated with oATP at 6 mmol/L or phosphate-buffered saline (PBS) alone daily for 4 weeks by i.p. injection and were sacrificed at the end of treatment by carbon dioxide inhalation. A group of age-matched wild-type (WT) C57Bl/6J male mice was used as internal control. The experiment was repeated in two separate 4-week trials, which both included 5 to 7 mice in each experimental group. Thus, the results are indicative of at least 10 animals per group. The Sgca mice were randomly assigned to the experimental groups, and the testing of samples was performed blindly (S.A. and S.Bal.).

Blood samples were collected by intracardial puncture at the end of the fourth week of treatment and centrifuged for 10 minutes at 7000 × g. Immediately after centrifugation, the serum was isolated and stored at −20°C. Serum creatine kinase levels were measured using the creatine kinase—N-acetyl cysteine quantitative system, according to manufacturer’s instructions (BPC BioSIED, Rome, Italy).

All animal experiments were performed in accordance with the Swiss Federal Veterinary Office guidelines and authorized by the Animal Studies Committee of Cantonal Veterinary.

Four-Limb Hanging Test

At the beginning of the study and at the end of the second and fourth week of treatment, the muscle strength of oATP-treated, PBS-treated Sgca, and WT control mice was scored through the four-limb hanging test. In this method, mice are subjected to a 180-second lasting hanging test, during which a falling score is recorded. In each of the two treatments, all of the mice had to hang for three trials, and the average maximum hanging time of the three trials was measured (standard operating procedure, http://www.treat-nmd.eu/research/preclinical/preclinical-efficacy-standards, last accessed December 1, 2017).

Reagents and Antibodies

oATP (Sigma-Aldrich, St. Louis, MO) was reconstituted at a final concentration of 100 mmol/L in PBS and stored at −80°C; the reconstituted drug was thawed and diluted in PBS at 6 mmol/L immediately before use. Lipopolysaccharide (LPS; Sigma-Aldrich) was reconstituted at a concentration of 1 mg/mL in Hanks’ balanced salt solution and stored at −20°C, ATP (Sigma-Aldrich) was reconstituted at 100 mmol/L (pH 7) in water with sodium hydroxide and stored at −20°C, and benzoyl ATP (BzATP; Sigma-Aldrich) was reconstituted at 10 mmol/L in water and stored at −20°C.

The antibodies and dilutions used in the study include the following: mouse monoclonal antibody to collagen type I [1:500 for Western blot (WB) analysis] and rat monoclonal antibody to Ly6C [1:500 for immunohistochemistry (IHC)] from Abcam (Cambridge, UK); rat monoclonal antibody to CD45 (1:10 for IHC) from BD Pharmigen (San Jose, CA); mouse monoclonal antibody to activating signal
cointegrator–1 (ASC-1) (1:500 for WB analysis) and rabbit polyclonal antibodies to glyceraldehyde phosphate dehydrogenase (1:500 for WB analysis) from Santa Cruz Biotechnology (Dallas, TX); rabbit polyclonal antibody to CD3 (1:20 for IHC); rabbit polyclonal antibodies to P2X7 receptor (R; extracellular) and P2X4R (1:50 and 1:300, respectively, for immunofluorescence) from Alomone Labs (Jerusalem, Israel); rabbit polyclonal antibody to transforming growth factor (TGF)–β (1:1000 for WB analysis) from Cell Signaling Technology (Danvers, MA); mouse monoclonal antibody to CD39 (1:1000) from Abcam; mouse monoclonal antibody to α-SG (1:100 for WB analysis) from Novocasta (Newcastle upon Tyne, UK); mouse monoclonal antibody to β-dystroglycan (1:50 for WB analysis) from Novocastar; mouse monoclonal anti–caveolin-3 (1:10,000 for WB analysis) from BD Bioscience (Milan, Italy); mouse monoclonal antibody to myosin heavy chain (fast; 1:1000 for WB analysis) from Novocastar; rabbit anti-mouse IgG horseradish peroxidase—conjugated (1:700 for WB analysis) antibodies from Dako (Glostrup, Denmark); donkey anti-rabbit IgG horseradish peroxidase—conjugated (1:5000 for WB analysis) antibodies from GE Health Care (Little Chalfont, UK); mouse monoclonal anti-rat κ and λ light chain horseradish peroxidase—conjugated (1:100 for IHC) antibody from Sigma-Aldrich; rat monoclonal to FOXP3 (1:20 for IHC) from ebioscience (Waltham, MA); and goat anti-mouse AlexaFluor 488 and donkey anti-rat AlexaFluor 594 (1:750 for immunofluorescence) antibodies from Thermo Fisher Scientific (Waltham, MA). The EnVision Detection System Peroxidase/Diaminobenzidine, rabbit and mouse, was from Dako. The Annexin A5-FITC/7-AAD Kit was purchased from Beckman Coulter (Marseille, France).

Satellite Cell Isolation from Total Muscle

Satellite cells were isolated from WT and Sgca mice at passage 8 to passage 10. Forelimb, hind limb, and diaphragm muscles were dissected, mechanically cut, and enzymatically digested at 37°C under constant shaking with a solution containing collagenase I (100 μg/mL; Sigma-Aldrich), dispase (500 μg/mL; Gibco, Waltham, MA), and DNaseI (100 μg/mL; Roche, Basel, Switzerland) in PBS (Sigma-Aldrich). Undigested tissue was precipitated for 5 minutes, and the supernatant was centrifuged for 5 minutes at 1200 × g. The cell pellet was resuspended in Dulbecco’s modified Eagle’s medium plus 10% fetal bovine serum, 1% L-glutamine, 1% penicillin/streptomycin, and 1% gentamicin; then, it was plated in 150-mm Petri uncoated dishes for 1 hour. After preplating, the nonadherent satellite cell—enriched population was collected and plated in collagen-coated (collagen from calf skin; Sigma-Aldrich) 90-mm Petri dishes at a density of 30,000 cells per Petri dish. After a few days in proliferation, the myoblasts were eventually plated at high density in differentiation medium (Dulbecco’s modified Eagle’s medium, 10% donor horse serum, 1% L-glutamine, 1% penicillin/streptomycin, 1% gentamicin, and 2.5 ng/mL basic fibroblast growth factor).

Histologic Studies

Sections (7 μm thick) from gastrocnemius (GN) and anterior tibialis muscles isolated from PBS-treated mice, oATP-Sgca mice, and WT controls were cut on a cryostat and stained with standard hematoxylin and eosin and acid phosphatase for inflammatory reactions and Masson trichrome stain.

Image analysis in terms of area fraction quantification, signal intensity, and object counting was performed using semi-automated measurement tools in NIS-Elements AR software version 4.20.00 (Laboratory Imaging, s.r.o., Prague, Czech Republic). Central nucleation (expressed as percentage of the total nuclei in each section) was quantified in a blinded manner (S.A. and S.Bal.) in three sections for each muscle (n = 10 mice for each experimental group, n = 400 to 600 myofibers per single animal) with the Axioplan Imager M2 microscope software AxioVs40 version 4.8.2.0 (Zeiss, Oberkochen, Germany). Paraffin-embedded diaphragms isolated from PBS-treated oATP-Sgca and WT control mice were cut in cross section at the midbelly of the muscle (7 μm thick). After deparaffinization, the slides were stained with hematoxylin and eosin12 and Picrosirius red staining for collagen fibers.

Immunohistochemistry

Cryosections (5 μm thick) of GN and anterior tibialis muscles isolated from PBS-treated, oATP-Sgca, and WT control mice were fixed in acetone at −20°C for 10 minutes, washed three times in PBS, and then washed twice in 0.025% PBS/Triton. Sections were incubated in 8% bovine serum albumin in PBS (blocking solution) for 1 hour at room temperature. Then, sections were incubated with primary antibodies diluted in 1% bovine serum albumin in PBS solution overnight at 4°C. The next day, sections were washed two times with 0.025% Triton in PBS, incubated 15 minutes in 0.3% H2O2 in PBS, and subsequently incubated with an Envision secondary antibody for 1 hour. After being rinsed two times with PBS (5 minutes each), the sections were stained with a diaminobenzidine substrate solution (freshly made just before use), counterstained with hematoxylin, washed with deionized water, dehydrated through five changes of alcohol (70%, 90%, 95%, 100%, and 100%) for 1 minute each, cleared in two changes of xylene (3 minutes each), and mounted with a Vectashield mounting medium (Vector Laboratories, Burlingame, CA). Slides were observed under a Leica Diaplan microscope (Leica, Wetzlar, Germany). The number of positive cells was blindly counted (S.Bar. and S.A.) on the whole sections per mouse. The total area of the section was measured and expressed in arbitrary units.

For immunofluorescence, unfixed cryosections (5 μm thick) of GN and anterior tibialis muscles were incubated in common antibody diluent (BioGenex, San Ramon, CA) for
10 minutes and after with primary antibodies (CD45, P2X4, and P2X7 antibodies) overnight at 4°C. After three washes with PBS, sections were incubated with fluorescent secondary antibodies in common antibody diluent for 1 hour at room temperature in the dark. After being washed three times with PBS, sections were mounted with Vectashield mounting medium with DAPI (Vector Laboratories). Images were obtained using a Leica SPE confocal microscope equipped with argon/He-Ne laser sources and an HCX PL APO CS 63.0/1.40 oil objective. During image acquisition, the 488 and 594 lasers were set at 20% energy, and the emission range was between 500 and 550 and between 650 and 700 nm for P2X 488 and CD45 594, respectively. The photomultiplier voltage gain was set to eliminate cell autofluorescence.

Single-plane images were taken at the center of cell thickness.

**Real-Time Quantitative PCR**

Total RNA was extracted from GN muscles (from 20 to 55 mg) isolated from PBS-treated mice, oATP-Sgca mice, and WT controls, according to the manufacturer’s protocol.
Quality and quantity of RNA were analyzed using a NanoDrop spectrophotometer (NanoDrop Technologies, Inc., Wilmington, DE). The cDNA was synthesized from 350 ng of total RNA with the iScript cDNA Synthesis Kit (Bio-Rad Laboratories, Hercules, CA). Each RNA sample was controlled for genomic DNA contamination without reverse transcriptase addition into a cDNA synthesis mixture.

For receptor-type tyrosine-protein phosphatase C (CD45), adhesion G-protein-coupled receptor E1 (ADGRE1), and early growth response protein 2 (EGR2) and adaptive [CD3, CD4, and T-cell surface glycoprotein CD8α chain (CD8A)] immunity, of Treg activation [forkhead box protein P3 (FOXP3), amphiregulin (AREG), and IL-10], and of proinflammatory cytokines [interferon IFN-γ (IFN-γ), IL-1β, and IL-6] was determined by real-time quantitative PCR (qPCR) was performed in triplicate with the 2^−DDCt method with two

![Figure 1](https://example.com/figure1.png)

**Figure 1**  Inflammatory response and T-regulatory cell (Treg) activation in Sgca mice. Total RNA isolated from the gastrocnemii of 10 Sgca and 7 age- and sex-matched WT mice was reverse transcribed. The pattern of expression of molecular markers of innate [receptor-type tyrosine-protein phosphatase C (PTPRC), adhesion G-protein-coupled receptor E1 (ADGRE1), and early growth response protein 2 (EGR2)] and adaptive [CD3, CD4, and T-cell surface glycoprotein CD8α chain (CD8A)] immunity, of Treg activation [forkhead box protein P3 (FOXP3), amphiregulin (AREG), and IL-10], and of proinflammatory cytokines [interferon IFN-γ (IFN-γ), IL-1β, and IL-6] was determined by real-time quantitative PCR. PPIA and RPL13A transcript levels were used as housekeeping genes. Data are expressed as medians ± first and third quartiles. PTPRC: WT = 0.11 (0.05 to 0.18), Sgca = 0.82 (0.39 to 1.05). ADGRE1: WT = 0.21 (0.09 to 0.24), Sgca = 0.85 (0.43 to 0.96). EGR2: WT = 0.05 (0.01 to 0.32), Sgca = 0.59 (0.29 to 1.02). CD3: WT = 0.25 (0.02 to 0.52), Sgca = 0.89 (0.06 to 1.36). CD4: WT = 0.06 (0.06 to 0.11), Sgca = 0.63 (0.44 to 1.05). CD8A: WT = 0.09 (0.04 to 0.17), Sgca = 0.06 (0.04 to 0.11). FOXP3: WT = 0.31 (0.30 to 0.31), Sgca = 0.43 (0.41 to 0.57). AREG: WT = 0.001 (0.001 to 0.17), Sgca = 0.40 (0.07 to 0.57). IL-10: WT = 0.02 (0.01 to 0.05), Sgca = 1.00 (0.39 to 1.18). IL-1β: WT = 0.02 (0.062 to 0.21), Sgca = 0.38 (0.31 to 1.05). IL-6: WT = 0.37 (0.22 to 0.489), Sgca = 0.67 (0.52 to 0.86). IFN-γ: WT = 0.06 (0.0 to 0.27), Sgca = 0.66 (0.49 to 1.24). Statistical analysis was performed using the U-test. *P < 0.05, **P < 0.01, and ***P < 0.001.

(DNase treatment). Quality and quantity of RNA were analyzed using a NanoDrop spectrophotometer (NanoDrop Technologies, Inc., Wilmington, DE).

The cDNA was synthesized from 350 ng of total RNA with the iScript cDNA Synthesis Kit (Bio-Rad Laboratories, Hercules, CA). Each RNA sample was controlled for genomic DNA contamination without reverse transcriptase addition into a cDNA synthesis mixture. For receptor-type tyrosine-protein phosphatase C (CD45), adhesion G-protein-coupled receptor E1 (F4/80), early growth response protein 2, CD3 ε chain, CD4, CD8α chain, interferon-γ (IFN-γ), IL-6, FOXP3, IL-10, amphiregulin, myogenin, osteopontin (secreted phosphorylated protein 1), connective tissue growth factor, and TGF-B1 transcript quantification, real-time quantitative PCR (qPCR) was performed in triplicate with the 2^−DDCt method with two
housekeeping genes (peptidylprolyl isomerase A and ribosomal protein L13a) by the Bio-Rad CFX Maestro software version 4.0.2325.0418. For each specific primer set, the efficiency was >95% and a single product was seen on the melting curve analysis.

For mouse P2X receptors and IL-1β quantification, qPCR was performed in triplicate with the 2^-ΔΔCT method. Total protein lysates that corresponded to 12.5 ng of total RNA, and premixed primer/probe sets were ordered from Applied Biosystems (Foster City, CA) (Table 2). Relative expression levels were normalized using the comparative Ct (ΔΔCT) method with the housekeeping gene Gapdh and actin-α1 (Acta1, a muscle specific gene, NM_001272041.1), by the Bio-Rad CFX Maestro software.

Western Blot Analysis

Total protein lysates were isolated from PBS-treated mice, oATP-Sgca mice, and WT GN muscles, as previously described. Equal amounts of protein (40 to 120 μg) were resolved in 8% to 15% SDS-PAGE gel and transferred onto polyvinylidene difluoride membranes (Immobilon PVDF, Billerica, MA). Membranes were blocked with 5% bovine serum albumin in PBS/0.1% Tween 20. Blots were then incubated overnight at 4°C with primary antibodies. Horseradish peroxidase anti-IgG was used to visualize bound primary antibodies with the chemiluminescence system (Bio-Rad Laboratories). Band intensities were evaluated by densitometry using the Alliance Mini HD9 AutoAutomatic system (Uvitec, Cambridge, UK).
Ecto-ATPase Activity

Satellite cells were seeded in 24-well plates (5.5 × 10^4 cells/well) and cultured until they completely differentiated into mature myotubes. After removal of the culture medium, cells were washed once with 1 mL Hanks’ balanced salt solution, and 0.35 mL Hanks’ balanced salt solution containing 0.3 mmol/L ATP was added. At various times (0, 5, and 15 minutes), 100-μl aliquots of the incubations were withdrawn and incubations were stopped by filtration with a multiscreen vacuum manifold using Immobilon-P membrane plates (Millipore, Bedford, MA). ATP degradation was determined by the phosphate high-performance liquid chromatography analysis, as previously described.13 Cells were lysed, and protein content in each well was determined by Bradford assay.

Apoptosis

Satellite cells were seeded in 24-well plates (6 × 10^4 cells/well) and cultured until they completely differentiated into mature myotubes. The cells were pretreated with LPS (1 μg/mL) for 4 hours and then incubated with 3 mmol/L ATP and 300 μmol/L BzATP for 16 hours. Then, myotubes were analyzed by flow cytometry, according to other studies that have adopted the same technique.14–16 Specifically, the cells were stained with the Annexin A5 FITC/7-AAD Kit, and apoptosis was evaluated by flow cytometry, according to the manufacturer’s instructions. Sample analysis was performed using a Gallios cytometer (Beckman-Coulter) and Kaluza software version 1.1 (Beckman-Coulter).

Statistical Analysis

Most of the analyzed variables were not normally distributed, as confirmed by the Shapiro-Wilk test, and therefore differences of quantitative parameters between two groups of mice were analyzed by the U-Mann Whitney test for nonparametric data. P < 0.05 was considered statistically significant. Data are expressed as medians ± first and third quartiles. P2X4: WT = 0.147 (0.140 to 0.249), Sgca = 0.907 (0.549 to 1.905). P2X7: WT = 0.217 (0.167 to 0.258), Sgca = 0.767 (0.362 to 1.630). Statistical analysis was performed using the U-test. *P < 0.05.

Results

Sgca Skeletal Muscle Displays an Increased Expression of Markers of Innate and Adaptive Immunity

To quantify the inflammatory response in muscles isolated from Sgca mice, the transcript levels of different immune cell markers and of distinct cytokines were analyzed, compared with muscles isolated from age-matched WT mice. α-SG—deficient muscle overexpressed leukocytes as well as macrophage surface proteins. Receptor-type tyrosine-protein phosphatase C (CD45), adhesion G-protein—coupled receptor E1 (F4/80), and early growth response protein 2

Figure 3  P2X4 and P2X7 purinergic receptors are significantly up-regulated in Sgca muscle tissue. Total RNA isolated from the gastrocnemii of 10 Sgca and 7 age- and sex-matched WT mice was reverse transcribed. The pattern of expression of P2X receptors (Rs) was determined by real-time quantitative PCR. P2X2R transcript levels were not measurable. P2X4R and P2X7R transcripts are significantly up-regulated in α-SG—deficient mice. Gapdh and Acta1 were used as housekeeping genes. Data are expressed as medians ± first and third quartiles. P2X4: WT = 0.147 (0.140 to 0.249), Sgca = 0.907 (0.549 to 1.905). P2X7: WT = 0.217 (0.167 to 0.258), Sgca = 0.767 (0.362 to 1.630). Statistical analysis was performed using the U-test. *P < 0.05.
were up-regulated by 7.5-, 4.1-, and 10-fold, respectively. CD3 transcripts were increased by 3-fold, with a 10-fold increase of the proportion of CD4-expressing lymphocytes, and a 30% reduction in CD8A mRNA levels. Moreover, CD4 induction was associated with a 1.8-fold up-regulation of FOXP3, a marker of CD4⁺ T-regulatory (Treg) cells.

The recruitment of immune cells in the muscle of Sgca mice was confirmed by a 4.7-, 1.8-, and 11-fold induction of the proinflammatory cytokines IL-1β, IFN-γ, and IL-6, respectively, but also by a 50- and 3-fold induction of the Treg-specific anti-inflammatory cytokine IL-10 and of the Treg-secreted growth factor amphiregulin, respectively (Figure 1).

**Sgca Primary Myotubes Are Characterized by a Defect in Ecto-ATPase Activity and Undergo Apoptosis on ATP Treatment**

Primary satellite cells were isolated from the forelimb, hind limb, and diaphragm muscles of passage 8 to passage 10 old Sgca and WT mice and cultured until they completely

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**Figure 4**  
P2X7R is overexpressed in Sgca muscle cells, whereas P2X4R is mainly expressed in CD45⁺ cells. **A:** Frozen muscle tissue sections of gastrocnemii (GN) from Sgca and WT mice were immunostained with a monoclonal anti-CD45 antibody (red) and a monoclonal P2X7R antibody (red). Representative images are shown. **White arrows** indicate examples of the positive cells for P2X7R. **B:** Frozen muscle tissue sections of GN from Sgca and WT mice were immunostained with a monoclonal anti-CD45 antibody (red) and a monoclonal P2X4R antibody (red). Representative images are shown. **Yellow arrows** indicate examples of the positive cells for P2X4R and CD45. $n = 4$ (A and B, Sgca and WT mice). Scale bar $= 25 \mu m$ (A and B). Original magnification, $\times 40$ (A and B).
For the evaluation of apoptosis, the cells were pretreated with 1 μg/mL LPS for 4 hours, incubated with 3 mmol/L ATP and 300 μmol/L BzATP for 16 hours, and then stained with the Annexin A5 FITC/7-AAD Kit. The feasibility of a citofluorimetric assay in myotubes has been previously determined. The number of apoptotic muscle cells was determined measuring the percentage of Annexin V+/7-AAD− and Annexin V+/7-AAD+ cells (early and late apoptosis, respectively). The pretreatment with LPS is supported by the observation that LPS caused a marked up-regulation of striatal P2X7 expression. ATP and, more potently, BzATP induced apoptosis, an effect that was further enhanced by a pretreatment with LPS (Figure 2D).

P2XR Is Expressed in Sgca Muscle Tissue

The pattern of expression of the P2XR was evaluated in muscle lysates from the GN of Sgca mice and age-matched WT controls. A specific up-regulation of P2X4R and P2X7R transcripts (sixfold and fourfold, respectively) was observed (Figure 3), whereas P2X2R transcript levels were not measurable (Figure 3). To determine whether P2X4R and P2X7R were expressed in myofibers or in infiltrating inflammatory cells, a coimmunostaining was completed with the pan-granulocyte marker C45 and P2X4 and P2X7R antibodies, respectively, in GN sections.

P2X4R was hardly detectable on the muscle cell membrane in WT mice, and it was up-regulated and colocalized with CD45-positive cells in the Sgca mutants (Figure 4A). P2X7R expression was weak in WT myofibers, whereas clear and specific P2X7R patches at the plasma membrane level were detected in Sgca muscle fibers, suggesting an induction and a higher functionality of the receptor because of its cell surface localization. This same feature has been previously shown in the monocytic subset of myeloid-derived suppressor cells isolated from tumor-bearing mice.

Targeting P2X Signaling in Vivo Improves Biochemical and Functional Parameters of Disease Progression in Sgca Mice

Given the representation of different P2XR in Sgca muscles, we aimed at initially achieving a pleiotropic antagonistic effect on eATP signaling cascade. Four-week–old Sgca mice were treated with oATP, a compound known to act on multiple P2X receptor molecules. oATP was administered via i.p. injections, at the dose of 6 mmol/L. Sgca mice injected with vehicle alone (PBS) and a group of age- and sex-matched WT C57Bl/6 mice served as controls. Mice were weighted at the beginning (baseline) of the treatment and then on a weekly basis. No significant difference was detected between the various groups (data not shown). The evaluation of muscle strength was completed by the four-limb hanging test at the beginning of the treatment (time 0) and at the end of the second and fourth weeks of treatment.

Figure 5 oATP improves muscle strength and ameliorates creatine kinase (CK) serum levels in Sgca mice. A: Four-limb hanging test was performed on PBS-Sgca (light gray line), oATP-Sgca (dark gray line), and age-matched WT controls (black line) at the beginning and at the end of the second and fourth week of treatment. B: Serum CK levels were evaluated at the end of the fourth week of treatment. Blood samples were obtained via cardiac puncture from PBS-Sgca, oATP-Sgca, and WT mice. Data are expressed as medians ± first and third quartiles. WT = 675 (130 to 1220), PBS-Sgca = 8460 (3835 to 24,635), oATP-Sgca = 5360 (3500 to 5440). Statistical analysis was performed using the U-test. Data are expressed as means ± SD (A). n = 12 animals (A); n = 10 animals for each experimental group (B). *P < 0.05 for oATP-treated mice versus PBS-treated mice.
As expected from the literature, the untreated Sgca group displayed a progressive worsening of muscle strength along time, whereas the oATP-Sgca cohort maintained the scores measured at time 0 (P < 0.05 for oATP-treated mice versus PBS-treated mice after 4 weeks) (Figure 5A).

Serum creatine kinase concentrations, a marker of muscle cell degeneration, were evaluated at the end of the trial by cardiac puncture. Creatine kinase was markedly increased in PBS-Sgca mice when compared with the WT cohort, whereas the treatment with the P2X antagonist led to a 35% decrease of this effect (Figure 5B).

**Inhibition of P2X Signaling Ameliorates Morphologic Features and Reduces the Amplitude of the Innate and Adaptive Immune Responses in Sgca Mice**

The fraction area and the intensity of the inflammatory reactions surrounding degenerative/necrotic muscle cells were evaluated in GN from PBS- and oATP-treated animals by acid phosphatase staining, which provides a red positive signal in activated macrophages and degenerative myofibers. These parameters were decreased in the oATP-treated group by 46% and 38%, respectively (P < 0.01 and P < 0.05, respectively) (Figure 6, A and B).

Muscle morphology was quantified in GN by standard hematoxylin and eosin staining. Central nucleation (percentage), a sign of dystrophic myopathic damage, was reduced by 14% in GN of oATP-treated animals (Figure 6C). Moreover, muscle tissue from untreated Sgca mutants, when compared with WT mice, displayed an increase of myogenin transcripts (Myog) compared with matched controls was reverse transcribed. Myogenin transcripts (Myog) were quantified by real-time quantitative PCR. Ppia and Rpl13a served as housekeeping genes. Data are expressed as medians ± first and third quartiles. WT = 0.30 (0.16 to 0.54), PBS-Sgca = 0.669 (0.19-1.39), oATP-Sgca = 0.217 (0.10 to 0.55). Statistical analysis was performed using the U-test.

Muscle morphology was quantified in GN by standard hematoxylin and eosin staining. Central nucleation (percentage), a sign of dystrophic myopathic damage, was reduced by 14% in GN of oATP-treated animals (Figure 6C). Moreover, muscle tissue from untreated Sgca mutants, when compared with WT mice, displayed an increase of myogenin transcripts (Myog) compared with matched controls was reverse transcribed. Myogenin transcripts (Myog) were quantified by real-time quantitative PCR. Ppia and Rpl13a served as housekeeping genes. Data are expressed as medians ± first and third quartiles. WT = 0.30 (0.16 to 0.54), PBS-Sgca = 0.669 (0.19-1.39), oATP-Sgca = 0.217 (0.10 to 0.55). Statistical analysis was performed using the U-test.

In diaphragms, P2X antagonism resulted in a 71% and 11% decrease in the areas and number of the inflammatory reactions surrounding the muscle cells in necrosis, respectively (hematoxylin and eosin staining). The beneficial effect of purinergic antagonism was associated also with a 16% reduction of central nuclei (P<0.05) (Supplemental Figure S1).

Immunostaining and measurement of the CD45+ and Ly6C+ leukocytes infiltrating the muscle tissue revealed a positive signal for both markers in different sites of the inflammatory responses surrounding necrotic fibers. When quantified by automated microscopy on the total section area, the number of CD45+ and Ly6C+ cells in the total

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**Figure 6**

oATP improves the extension of reactive inflammatory infiltrates and morphologic features in Sgca mice. A: Frozen muscle tissue sections of gastrocnemii (GN) from PBS-Sgca and oATP-Sgca mice were collected at the fourth week of treatment and stained with acid phosphatase. **Black arrows** indicate inflammatory infiltrates. Representative images are shown. B: Graph of the analysis of the fraction areas and intensities of inflammatory infiltrates/total section area of muscles represented in A. Data are expressed as medians ± first and third quartiles. Area: WT = 0.006 (0.002 to 0.0267), PBS-Sgca = 0.117 (0.09 to 0.14), oATP-Sgca = 0.063 (0.04 to 0.09). Signal intensity: WT = 71.886 (12,400 to 217,000), PBS-Sgca = 6.30 × 10^6 (4.01 × 10^6 to 9.12 × 10^6), oATP-Sgca = 3.85 × 10^6 (2.53 × 10^6 to 6.13 × 10^6). Statistical analysis was performed using the U-test. C: The number of central nuclei was quantified in 600 fibers of three consecutive fields for each muscle section stained with hematoxylin and eosin. Data are expressed as medians ± first and third quartiles. WT = 2.00 (1.00 to 3.00), PBS-Sgca = 82.91 (80.18 to 89.34), oATP-Sgca = 70.89 (64.76 to 75.01). Statistical analysis was performed using the U-test. D: Total RNA isolated from GN lysates from PBS-Sgca, oATP-Sgca, and WT age-matched controls was reverse transcribed. Myogenin transcripts (Myog) were quantified by real-time quantitative PCR. Ppia and Rpl13a served as housekeeping genes. Data are expressed as medians ± first and third quartiles. WT = 0.054 (0.037 to 0.076), PBS-Sgca = 0.669 (0.19-1.39), oATP-Sgca = 0.217 (0.10 to 0.55). Statistical analysis was performed using the U-test. n = 10 (A, PBS-Sgca and oATP-Sgca mice, C, mice for each experimental group, and D, PBS-Sgca, oATP-Sgca, and WT mice). *P < 0.05, **P < 0.01. Original magnification, ×10 (A). AU, arbitrary unit.
areas was inhibited by 45% and 25%, respectively, in oATP-treated mice (Figure 7, A and B).

Accordingly, the protein levels of ASC-1, a coactivator of NF-κB, were undetectable in WT mice, markedly increased in Sgca muscle lysates, and significantly inhibited after P2X blockade (Figure 7C).

The immunostaining and measurement of the CD3 lymphocytes infiltrating the muscle tissue showed a CD3-positive signal in different sites of reactive inflammatory responses. When quantified by automated microscopy on the total section area, the fraction areas of CD3-positive signal were reduced by 52% in oATP-treated mice (data not shown).

Despite the decrease in the total number of CD3-positive cells, FOXP3⁺ cells were increased in muscle tissue of oATP-treated Sgca mutants by 34% (Figure 8, A and B).

The mechanisms of how oATP affects innate and adaptive response in Sgca skeletal muscle were also investigated by quantification of muscular IL-1β, IL-6, and IFN-γ transcripts. When compared with the PBS cohort, these cytokines were reduced by 49%, 39%, and 44%, respectively, in the oATP group (Figure 8C).

**Targeting P2X Signaling in Vivo Decreases Muscular Fibrotic Degeneration, Reducing the Transcription of Muscle Profibrotic Factors in Sgca Mice**

In accordance with this anti-inflammatory image, P2X blockade led to a reduction of the transcription of fibrogenic factors, which ultimately stimulate endomysial fibrosis and connective replacement of muscle tissue. The transcripts of
osteopontin (secreted phosphorylated protein 1), connective tissue growth factor, and TGF-β were quantified in total muscle lysates isolated from the GN of WT, PBS-treated, and oATP-treated Sgca mice and displayed a 79%, 45%, and 77% reduction in the presence of eATP blockade, respectively (Figure 9).

The effects of oATP on collagen type I and on TGF-β were also confirmed at the protein levels. P2X antagonism prevented the up-regulation of collagen type I observed in the untreated cohort, and it led to a reduction of both dimeric (25 kDa) and monomeric (15 kDa) active TGF-β isoforms (Figure 10A).

Accordingly, histologic analyses by Masson trichrome staining on GN sections revealed that oATP treatment led to a 20% decrease of the fraction area (Figure 10, B and C). Staining with Picrosirius red in diaphragms corroborated a 40% and 43% decrease of the fraction area and the intensity of fibrotic tissue in the oATP-treated group, respectively (Supplemental Figure S2).

**Discussion**

In sarcoglycanopathies, the mechanically weaker plasma membrane is easily damaged during muscle contraction, allowing release of intracellular antigens, infiltration of immune cells, and induction of profibrotic cytokines and growth factors. Although different experimental models...
have shown how distinct anti-inflammatory approaches may enhance stem cell therapy in mice. The molecular mechanisms that trigger the immune-mediated damage in this disorder have not been described yet. Likewise, limited are the clinical studies and the clinical experience on anti-inflammatory strategies in human sarcoglycanopathies.

Conversely, in Duchenne muscular dystrophy, it has been extensively highlighted that the immune system exerts a dichotomous role because two types of inflammation that promote and repair muscle injury are activated in the muscle tissue. Type 1 inflammation is characterized by increased expression of IFN-\(\gamma\) and is counterregulated by type 2 cytokines, such as IL-4 and IL-13, and the Treg anti-inflammatory cytokine IL-10.

Herein, it was found that type 1 and type 2 inflammation are activated also in \(\alpha\)-SG deficiency. When compared with WT animals, \(\alpha\)-SG–deficient muscle tissue displayed higher expression levels of proinflammatory cytokines, such as IL-1\(\beta\), IFN-\(\gamma\), and IL-6, augmented content of the proinflammatory molecule ASC-1, and increased infiltration of CD45 leukocytes, macrophages (innate immunity), and CD4\(^+\) lymphocytes (adaptive immunity). This process was counteracted by an increase in the number of FOXP3\(^+\) CD4\(^+\) Tregs, which also showed an activated phenotype, as confirmed by induced levels of the cytokine IL-10.

The potential role of the DAMP molecule eATP was next studied in the development of this tissue-specific immune-mediated damage, and it was shown, for the first time, that P2X-purinergic signaling is involved in this pathologic cascade.

Sgca mice were characterized by an enhanced expression of P2X4 and P2X7 receptors in muscle tissue, confirming that SG defects, as dystrophinopathies, result into a purinergic pathway overactivation. Although P2X4R was mainly up-regulated in CD45\(^+\) inflammatory cells infiltrating the muscle, P2X7R was overexpressed on the plasma membrane of Sgca muscle fibers. Noteworthy, in dysrophic cells, the receptor molecules were found to be organized in specific patches, suggesting in the monocyte subset of myeloid-derived suppressor cells isolated from tumor-bearing mice.

An excess of ATP in the extracellular milieu can be counteracted by the action of the ecto-ATPases CD39 and CD73, which in subsequent order convert ATP to adenosine. However, also, \(\alpha\)-SG displays in its C-terminal domain an ecto-ATPase function, and its absence causes a more consistent accumulation of ATP in the culture medium of primary myotubes. In turn, as already shown in distinct cell systems, an excessive extracellular ATP stimulation is able to exert a direct toxic effect in muscle cells because it induces apoptosis. The molecular basis of the proapoptotic effect can be because of an effect on mitochondrial metabolism or the association with pannexin molecules and the formation of a cytolytic pore (ie, a large conductance channel).

According to these first observations, the pharmacologic inhibition of purinergic signaling via oATP led to an improvement of muscular function and structure, a reduction of the innate/adaptive immune response and fibrosis, and an increase in FOXP3\(^+\) CD4\(^+\) Treg muscle infiltration. In this scenario, it is likely that the oATP-mediated boosting of Treg muscle infiltration played a role in ameliorating the Sgca mice phenotype. It has been shown that skeletal muscle resident Treg, producing IL-10 and expressing Areg, controls the switch of type 1 to type 2 inflammation in injured muscle and acts directly on muscle satellite cells, improving muscle repair.
isolated from mdx mice showed an increased cytosolic Ca\(^{2+}\) ion influx and IL-1\(\beta\) release when stimulated with BzATP,\(^{31}\) thus evoking an active role of skeletal muscle cells in the inflammatory mechanism through P2X-purinergic signaling. The assessment of Sgca and LGMD2D primary myoblast behavior under eATP stimulation and the direct effect of P2XR inhibition on these cells will be evaluated in our future studies.

\(\alpha\)ATP was originally described as an irreversible P2X7R antagonist\(^{32}\) but was shown later to also block other P2XRs, such as the P2X4 subtype,\(^{33}\) making this compound a good candidate for a first proof of principle of our hypothesis in Sgca mice, in which an up-regulation of both P2X7 and P2X4 receptors was found. \(\alpha\)ATP has also demonstrated promise as a treatment modality in several models of autoimmune diseases. Successful examples are experimental models of collagen-induced arthritis, autoimmune diabetes, allergic encephalitis, autoimmune colitis, and allograft rejection in cardiac and islet transplantation.\(^{33-39}\) In these disorders, the compound works as a suppressor of the innate and adaptive immunity as well as an inducer of Treg expansion.

However, numerous other P2XR antagonists, more efficient and more specific, have been discovered in the past few years, especially anti-P2X7R.\(^{40-43}\) CE-224,535 has been tested in clinical trials for rheumatoid arthritis in patients with an inadequate response to methotrexate. The drug was not efficacious, compared with placebo, but demonstrated an acceptable safety and tolerability profile (NCT00628095).\(^{44,45}\) The purinergic P2X7 antagonist AZD9056 was shown to have the potential to improve symptoms in patients with moderate-to-severe Crohn disease combined with a beneficial risk profile. AZD9056 was well tolerated, and no serious adverse events were reported.\(^{46}\) The molecule GS1482160 has already been explored as a possible tool to detect neuroinflammation, and a phase I clinical study in humans is currently undergoing (NCT00849134).\(^{47}\)

Currently, no therapies are available for LGMD2D patients. Many studies in each of the four sarcoglycan deficiencies demonstrated that viral-mediated gene transfer, using adeno-associated vectors, can correct the pathologic signs of the disease in animal models.\(^{48-56}\) However, some previous \(\alpha\)-SG gene transfer experiments revealed the

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**Figure 10** \(\alpha\)ATP induces a decrease of collagen profibrotic factors in muscles from Sgca mice. A: Total protein lysates, isolated from gastrocnemii (GN) of PBS-Sgca, \(\alpha\)ATP-Sgca, and WT age-matched controls, were subjected to Western blot analysis. Membranes were probed with a rabbit polyclonal anti-collagen type I and a mouse monoclonal anti—transforming growth factor-\(\beta\) (TGF-\(\beta\); the 25-kDa dimer and the 15-kDa monomer) antibody. Representative images are shown. B: Frozen muscle tissue sections of GN from PBS-Sgca, \(\alpha\)ATP-Sgca, and WT age-matched controls were collected at the fourth week of treatment and stained with a standard Masson trichrome stain protocol. Representative images are shown. C: Graph of the fraction areas of fibrotic blue positive signal/arbitrary unit (AU) of total section area of muscles represented in B. Data are expressed as medians ± first and third quartiles. WT = 0.029 (0.026-0.031), PBS-Sgca = 0.077 (0.076 to 0.107), \(\alpha\)ATP-Sgca = 0.061 (0.052 to 0.067). Statistical analysis was performed using the \(U\)-test. \(n = 6\) (A, PBS-Sgca, \(\alpha\)ATP-Sgca, and WT mice); \(n = 5\) (B, PBS-Sgca, \(\alpha\)ATP-Sgca, and WT mice). \(*P < 0.05\) for WT versus Sgca. Original magnification, \(\times 10\) (B). GAPDH, glyceraldehyde-3-phosphate dehydrogenase.
occurrence of specific immune response, leading to generation of α-SG—specific antibody, T-cell infiltration of treated muscles, and disappearance of the vector. In this respect, we suggest that P2X-purinergic signal antagonism, with its double anti-inflammatory effect (inhibition of innate/adaptive inflammatory response and enhancement of immune tolerance by Treg expansion), could be associated with gene transfer therapies for the treatment of LGMD2D to obtain a more stable α-SG expression.

Moreover, this study is in agreement with recent lines of research, which emphasize the relevance of DAMPs in chronic degenerative disorders. Intriguingly, these sophisticated molecules are danger signals important not only for the inflammatory response but also for tissue repair because they can orchestrate tissue healing by acting on different receptors. Thus, better understanding of the consequences of DAMP signaling on muscle inflammation and on muscle regeneration is the key to promote tissue repair and healing.

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Supplemental Data

Supplemental material for this article can be found at https://doi.org/10.1016/j.ajpath.2018.10.008.

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