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A crucial physiological role of Piezo1 channel in differentiation rather than proliferation during myogenesis

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EDITORIAL**A crucial physiological role of Piezo1 channel in differentiation rather than proliferation during myogenesis**Ming Jie Wang¹, Yi Chun Zhu*¹, Jian Shi*^{1,2}

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Mechanical signals are critical for the growth, maintenance and repair of skeletal muscle. However, it remained elusive whether **the mechanically activated ion channels, notably** the novel Piezo1 channel, mediate these functions in the muscle. **Although an earlier report¹ suggested that Piezo1 channel could be important for myoblast cell fusion and myotube elongation in the C2C12 cell line, the current study by Bosutti et al² made significant progress in determining the physiological roles of Piezo1 channel in myogenesis and adult myofibre activity. The novelty presented by this paper exists in three aspects. Firstly, the functional roles of Piezo1 channel were thoroughly investigated in primary cells and skeletal muscle under physiological condition for the first time. Not only the differentiation, but also the proliferation and contraction were analysed against Piezo1 channel in this paper. Secondly, a precise**

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3 time-window for intervening in the myogenic differentiation was identified, generating
4 new understanding of how temporal modulation of Piezo1 would promote
5 myogenesis. Thirdly, the chemical activator of Piezo1 channel, Yoda1, was proved to
6 be effective in inducing differentiation without causing any obvious side effect after
7 the long-term treatment, which provides the direct evidence that a novel
8 pharmacological approach to skeletal muscle regeneration is achievable. These
9 findings raise important questions in terms of functional significance of Piezo1
10 channel and its translational potential.

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12 Mechanical signals generated from cellular environment initiate and promote
13 myogenesis during which satellite cells (SC cells), the muscle precursor cells, are
14 directed to proliferate, differentiate and migrate with temporal and spatial patterns.
15 The scale and length of these mechanical signals can change dynamically,
16 depending on the reciprocal interaction between the cells and their niche³. For this
17 reason, the mechanosensors which translate the mechanical signals into biological
18 signals are indispensable for the determination of the cell fate at any given time. As a
19 relatively newly discovered mechanosensor, Piezo1 channel is a bona fide sensor of
20 mechanical force in mammals⁴, which has been unanimously agreed by independent
21 groups across the world. Named after the Greek word for pressure, Piezo1
22 assembles as a trimer to form Ca²⁺- and Na⁺-permeable ion channels with a typical
23 unitary conductance of ~25 pS⁴. Mouse Piezo1 consists of 2547 amino acids, while
24 human Piezo1 is composed of 2521 amino acids. Physiological studies have
25 previously revealed that Piezo1 channel contributes to the blood pressure regulation,
26 exercise performance facilitation, vascular development, red blood cell volume
27 sensation, baroreception, cartilage stress sensation, bone growth and
28 osteogenesis⁴. But it still remains underexplored whether and how Piezo1 channel
29 determined myogenesis and myofibre activities at physiological levels. The research
30 on Piezo1 functional roles in these tissue or cells was also complicated by the co-
31 expression of other mechanosensitive ion channels such as TRP channels, as all
32 these channels could be simultaneously activated by a mechanical stimulus. Thus
33 identification of a chemical agonist specific for Piezo1 channel would make the
34 distinguishment possible. Screened from ~3.25 million compounds with an unbiased
35 cell-based fluorescence assay, Yoda1 with its specificity against Piezo1 has been
36 widely tested⁴. Many studies have convincingly demonstrated that Yoda1 is highly
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3 selective for Piezo1 channel. Even Piezo2 channel, the only paralog of Piezo1, is
4 completely insensitive to Yoda1. The research on the vasculature confirmed that
5 Yoda1 could even mimic the effects of laminar shear stress (mechanical force) on
6 both the physiological and pathophysiological functions of endothelial cells^{4,5}.
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8 Therefore Yoda1 turns out to be a powerful tool for the functional investigation into
9 Piezo1 channel.
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14 In the current study, the SC cells characterised with Pax-7, an **exclusively** expressed
15 transcription factor in these cells, were chosen for the dissection of Piezo1
16 **physiological functions**. **Neither** the total cell number **nor** the percentage of Pax-7-
17 positive cells **was** changed by the **treatment** of Yoda 1 at **a concentration** which
18 would activate most Piezo1 channels. In contrast, the SC cell differentiation was
19 triggered by the same concentrations of Yoda1 as hallmarked by the morphology of
20 **cell** elongation and visualization of myogenin (MyoG), the muscle-specific
21 transcription factor. It is hence concluded that Piezo1 channel is only crucial for SC
22 cell differentiation without playing any noticeable role in proliferation and apoptosis.
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26 The further analysis of “time window” showed that the differentiation only occurred
27 after the cells were treated with Yoda1 for 72 hours, indicating that sophisticated
28 downstream events took place following the activation of Piezo1 channels during this
29 process. Interestingly, the differentiation **was** accompanied with a highly-regulated
30 alignment of SC cells as most cells were oriented towards the same direction. The
31 alignment was also clearly observed in endothelial cells following the activation of
32 Piezo1 channels by mechanical force⁵, which implies that a general mechanism may
33 underlie the Piezo1-guided alignment in all cell types. Future in-depth mechanistic
34 research will determine **how such spatial and temporal control is achieved in Piezo1-**
35 **mediated differentiation of muscle precursor cells**.
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40 After the differentiation of muscle precursor cells into myofibres, **it is also completely**
41 **unknown whether Piezo1 channels regulate the myofibre physiology**. In this paper,
42 the authors measured the electrical activity and contractility before and after the
43 activation of Piezo1 by Yoda1 in the adult muscle fibres. For the electrical activity,
44 neither the amplitude of the miniature endplate current (MEPCs) nor the frequency of
45 endplate current (fMEPCs) was altered by Yoda1 after the stimulation of the
46 neuromuscular junction (NMJ) synapse. For the contractility, the muscle contraction
47 following an electrical stimulation to the motor nerve was also not changed by
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3 Yoda1. Further Ca^{2+} imaging experiments showed that Piezo1-mediated Ca^{2+} entry
4 was nearly lost in adult skeletal muscle fibres. Only some sporadic Ca^{2+} events from
5 Piezo1 channels could still be observed when a higher concentration of Yoda 1 was
6 used. As positive control, other Ca^{2+} signals were normal and expected. So these
7 novel results rule out a physiological role of Piezo1 in these investigated functions of
8 adult myofibres. **However, these findings should also be interpreted with caution as it**
9 **is still possible for Piezo1 to play a role in other functions of myofibres such as**
10 **chronic adaptation to exercise.**

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12 As complementary approaches, **the methods of** biochemical immunostaining and
13 single-cell sequencing were adopted to validate the functional results of Piezo1
14 channels in myogenesis and myofibre **physiology**. To reduce any potential concern
15 about the Piezo1 antibody specificity, the authors performed the antigen retrieval
16 control experiments with blocking peptide which appear to be convincing. It was then
17 found that Piezo1 expression **level** was gradually decreased when the differentiation
18 progressed. When the differentiation of muscle precursor cells was completed and
19 mature myofibres were formed, Piezo1 expression seemed to be the lowest. In
20 mature myofibres, the Piezo1 channels were only located to a small endplate region
21 where the channels also had the least Ca^{2+} permeability, which is consistent with the
22 findings that the channels are not required for the investigated physiological
23 functions of myofibres. **The interesting expression pattern of Piezo1 during**
24 **myogenesis reported in this paper makes one wonder how Piezo1 channels are**
25 **expressed in myofibre dedifferentiation and what their functional roles are in that**
26 **scenario. Follow-up studies will certainly be needed to answer these intriguing**
27 **questions.**

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29 The current study reveals for the first time that the novel **mechanically gated** ion
30 channel, Piezo1, determines the differentiation **during** myogenesis **in a temporal**
31 **manner**. On the contrary, the channel **doesn't play any physiological role in other**
32 **functions** such as proliferation during myogenesis (as shown in Figure 1). **As Piezo1**
33 **activation can initiate the differentiation in muscle precursor cells**, decoding the
34 downstream cellular mechanism will potentially generate even deeper insight into the
35 understanding of mechanical force in muscle physiology⁶.

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37 **As mouse Piezo1 and human Piezo1 share high similarity at both protein structure**
38 **and function**, the knowledge arising from this study can be readily extended to any
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3 future research which investigates Piezo1 channelopathies in human myogenesis
4 abnormalities. **For such investigation**, sample genetic analysis from the patients
5 suffering from these disorders will likely generate some important clinical information.
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7 In addition, the current study may open a new avenue for the treatment of skeletal
8 muscle injury or progressive myopathies. **Because the differentiation initiated by**
9 **Piezo1 channels could be critical to muscle regeneration, fine-tuning of Piezo1**
10 **channel activity with pharmacological approaches will probably bring therapeutic**
11 **benefits for the recovery.**
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17 Therefore, pharmacological modulation of Piezo1 channel activity could represent a
18 novel strategy to **regulate** physiological or pathophysiological functions. The
19 modulators of Piezo1 channel discovered **so far** include the agonist Yoda1 (its
20 derivatives as well) and some lipid molecules^{1, 7}. They make it highly likely to
21 **modulate** the channel's activation (opening) or inactivation (closure) at a designated
22 time. From this perspective, these modulators or their derivatives could have
23 tremendous potential to be developed into therapeutic drugs in future for the
24 treatment of some myogenic diseases or **other Piezo1 channelopathies.**
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32 In conclusion, the important findings from this study highlight a novel physiological
33 role of Piezo1 channel in skeletal myogenesis. The new information about the
34 specific time window also sheds light on a potential therapeutic strategy to regulate
35 skeletal muscle physiology and pathophysiology.
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51 **CONFLICT OF INTEREST**

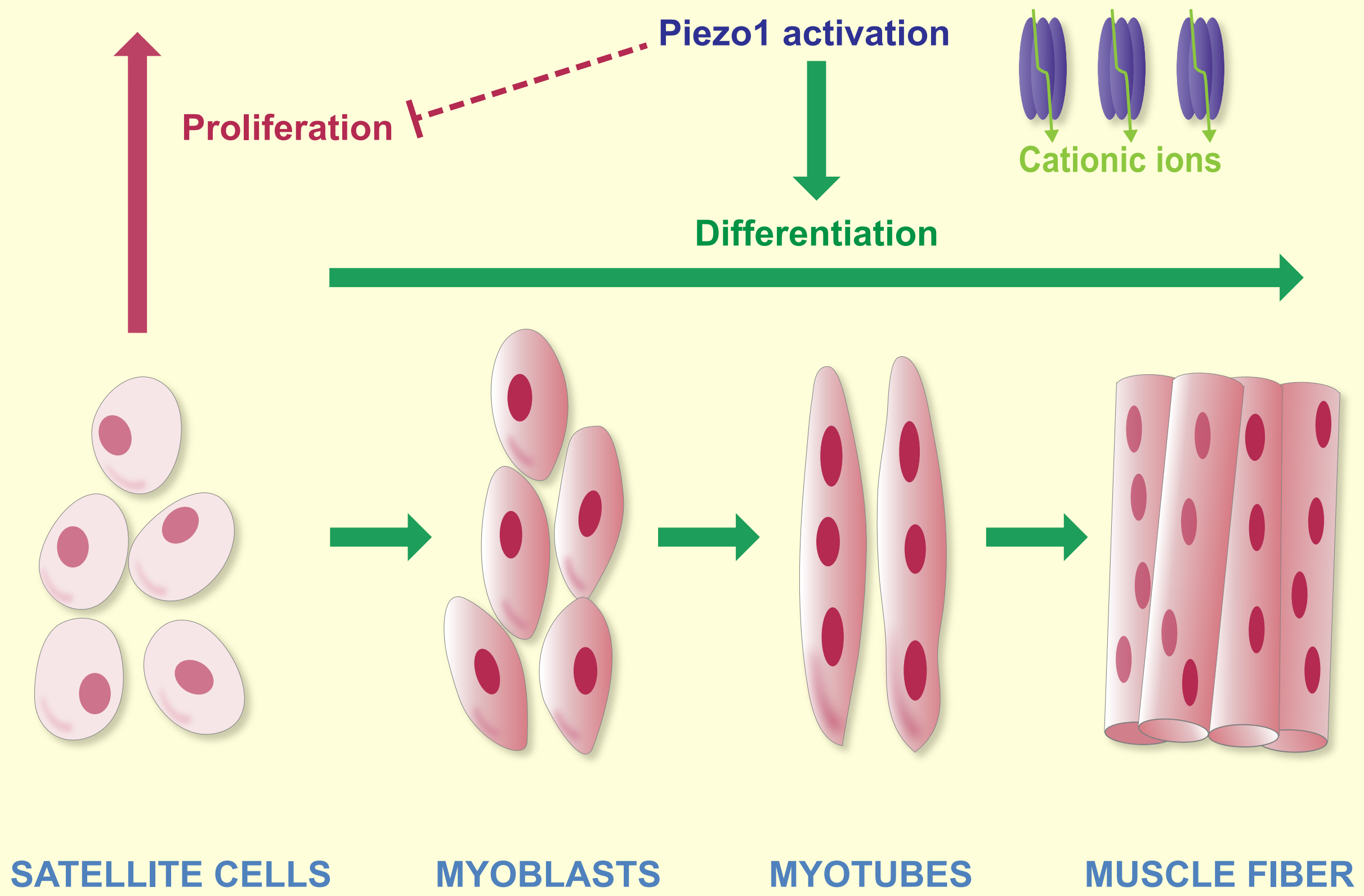
52 The authors declare no conflict of interest.
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SATELLITE CELLS

MYOBLASTS

MYOTUBES

MUSCLE FIBER