Maurocalcin phosphorylated at threonin 26 maintains its activity on ryanodine receptor-mediated $\text{Ca}^{2+}$ release in intact muscle fibers

Dóra Bodnár*, Laszlo Csernoch*, and Vincent Jacquemond*1

Ronjat et al. (1) report that a phosphorylated form of the scorpion venom toxin maurocalcin (MCa) loses its ability to activate its preferred target, the type 1 ryanodine receptor (RYR1). RYR1 is the sarcoplasmic reticulum (SR) $\text{Ca}^{2+}$ release channel responsible for the rise in cytosolic $\text{Ca}^{2+}$, which activates contraction during excitation–contraction coupling of skeletal muscle. Ronjat et al. report that phosphorylation of MCa threonine 26 (MCaThr26) completely reprograms the toxin, converting it into a RYR1 blocker. To assess the function of MCa and of MCaThr26-Phospho, the authors relied on several in vitro assays, including measurements of current through RYR1 channels reconstituted into lipid bilayer and $[^3H]$-ryanodine binding and $\text{Ca}^{2+}$ release from SR vesicles. However, Ronjat et al. did not obtain data from intact muscle fibers where RYR1 functions in its native physiological environment, being activated by its molecular partner, the voltage-sensing protein CAV1.1 in the plasma membrane. This is unfortunate because MCa has a unique effect on RYR1-mediated SR $\text{Ca}^{2+}$ release in intact muscle fibers, making this assay both reliable and physiologically relevant: indeed, under these conditions, MCa transiently keeps RYR1 channels from closing following membrane repolarization. This process manifests in a remarkable prolongation of the voltage-activated cytosolic $\text{Ca}^{2+}$ transient (2, 3) after the end of voltage activation.

We had the chance to test the effect of MCaThr26-Phospho, together with MCaThr26-E, where threonin is replaced with a glutamic acid residue, in intact muscle fibers. We believe that the results are interesting in regard to the issue raised. Fig. 1A shows examples of voltage-clamp activated cytosolic $\text{Ca}^{2+}$ transients in muscle fibers equilibrated with either MCa (10 μM) or MCaThr26-Phospho (10 μM) or MCaThr26-E (100 μM) in the voltage-clamp pipette. Experiments were conducted under conditions routinely used in our laboratory (4). For each fiber, we show the change in rhod-2 fluorescence in response to a 10- and 100-ms-long pulse, from −80 to +20 mV. As reported previously (2), the effect of MCa is use-dependent (i.e., the longer the depolarization, the larger the effect). Thus, the 100-ms pulse in the presence of MCa generates a prominent, sustained elevation of cytosolic $\text{Ca}^{2+}$ after the pulse. The exact same effect was observed with MCaThr26-Phospho but not with MCaThr26-E. Fig. 1B shows mean values for the peak (Fig. 1B, Upper) and for the end of record (~11 s following repolarization) (Fig. 1B, Lower) $\text{Ca}^{2+}$ levels, in the different conditions. There was no effect of MCa or of either derivative on the peak; however, $\text{Ca}^{2+}$ levels remained elevated after repolarization in a use-dependent manner for both MCa and MCaThr26-Phospho.

Altogether, we believe that the above results are an important complement to the data presented by Ronjat et al. (1) and there are reasons worth discussing as to why MCaThr26-Phospho becomes inefficient in vitro assays but remains active in intact muscle fibers. One possibility could be that high phosphatase activity inside the muscle fibers so completely dephosphorylates intracellular MCaThr26-Phospho as to release active MCa. If so, one may reasonably doubt that intracellular PKA activity would have the potential to inactivate native MCa in the fibers.

Acknowledgments

We thank Michel De Waard for the generous gift of MCa, MCaThr26-Phospho, and MCa Thr26-E. This work was supported by grants from the Centre National de la Recherche Scientifique, the Institut National de la Santé et de la Recherche Médicale, and the Université Claude Bernard Lyon 1 to the Institut NeuroMyoGène, and by the French-Hungarian Hubert Curien Partnership program Balaton.

*Department of Physiology, Faculty of Medicine, University of Debrecen, Debrecen, Hungary 4032, and †Institut NeuroMyoGène, CNRS UMR 5310, INSERM U1217, Université Claude Bernard Lyon 1, Villeurbanne 69622, France

Author contributions: V.J. designed research; D.B. and V.J. performed research; D.B., L.C., and V.J. analyzed data; and L.C. and V.J. wrote the paper.

The authors declare no conflict of interest.

1To whom correspondence should be addressed. Email: vincent.jacquemond@univ-lyon1.fr.

Fig. 1. MCaThr\textsuperscript{26}-Phospho is as active as MCa on voltage-activated SR Ca\textsuperscript{2+} release. (A) Changes in rhod-2 fluorescence (F/F\textsubscript{0}) elicited in muscle fibers equilibrated with (from top to bottom) the control intracellular solution, the intracellular solution with 10 μM MCa, 10 μM of MCaThr\textsuperscript{26}-Phospho, and 100 μM of MCaThr\textsuperscript{26}-E. Each fiber was stimulated by a 10- and then a 100-ms-long voltage-clamp depolarization. The inset shows the beginning of the records in the control fiber on an expanded time-scale. (B) Mean values for the change in rhod-2 fluorescence (ΔF/F\textsubscript{0}) for the initial peak change (Upper) and for the end of record level (Lower) in the different conditions (control, n = 7; MCa, n = 5; MCaThr\textsuperscript{26}-Phospho, n = 5; MCaThr\textsuperscript{26}-E, n = 6). Data values are presented as means ± SEM for n fibers. Statistical significance was determined using a Student’s t test (*P ≤ 0.05, **P ≤ 0.01).