## **RSC** Advances



## CORRECTION



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## Correction: Rapid synthesis of internal peptidyl $\alpha$ -ketoamides by on resin oxidation for the construction of rhomboid protease inhibitors

Tim Van Kersavond,<sup>a</sup> Raphael Konopatzki,<sup>a</sup> Merel A. T. van der Plassche,<sup>b</sup> Jian Yang<sup>b</sup> and Steven H. L. Verhelst<sup>\*ab</sup>

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Correction for 'Rapid synthesis of internal peptidyl  $\alpha$ -ketoamides by on resin oxidation for the construction of rhomboid protease inhibitors' by Tim Van Kersavond *et al.*, *RSC Adv.*, 2021, **11**, 4196–4199, DOI: 10.1039/D0RA10614C.

The authors regret that an incorrect version of Fig. 1 was presented in the original manuscript. The R-group in compound 7 was incorrectly indicated as  $(CH_2)_5$ Ph. The corrected version of the figure with the R group as  $(CH_2)_4$ Ph is shown below. The Royal Society of Chemistry apologises for these errors and any consequent inconvenience to authors and readers.

<sup>a</sup>Leibniz Institute for Analytical Sciences ISAS, e.V., Otto-Hahn-Str. 6b, 44227 Dortmund, Germany

<sup>b</sup>KU Leuven, Department of Cellular and Molecular Medicine, Laboratory of Chemical Biology, Herestr. 49 box 802, 3000 Leuven, Belgium. E-mail: steven.verhelst@kuleuven.be



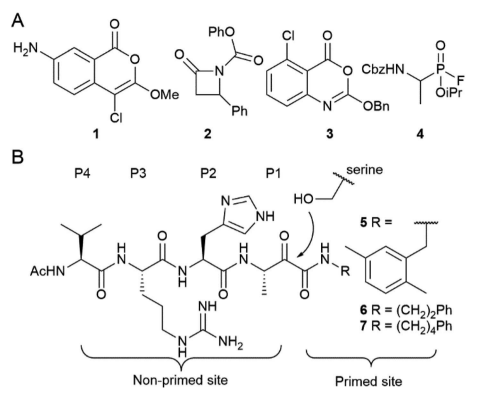


Fig. 1 Examples of rhomboid inhibitors. (A) 4-Chloro-isocoumarins (1),  $\beta$ -lactams (2), benzoxazinones (3) and fluorophosphonates (4). (B)  $\alpha$ -Ketoamide rhomboid inhibitors (5–7). The peptidic element in the non-primed site is indicated with the P1–P4 position according to the Schechter and Berger protease substrate nomenclature.<sup>1</sup>

## References

1 I. Schechter and A. Berger, Biochem. Biophys. Res. Commun., 1967, 27, 157-162.