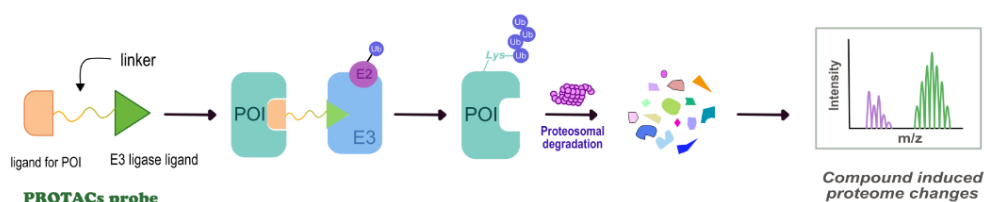


PROTACs as a target deconvolution tool of Hedgehog Pathway Inhibitor 1M. Bagka¹, S. Hoogendoorn^{1*}¹University of Geneva, Department of Organic Chemistry

The Hedgehog (Hh) signaling pathway is pivotal for embryonic development of vertebrates, however abnormal activation can lead to tumorigenesis. The current FDA approved drugs targeting the Hh pathway act on the upstream protein Smoothened and suffer from acquired resistance, rendering essential to find new inhibitors with a different target¹. Hedgehog Pathway Inhibitor 1 (HPI-1) was reported as a downstream inhibitor by Hyman et al and has subsequently been shown to have anti-cancer activity, but its cellular target has remained elusive for many years².

Target identification by protein degradation and label-free quantitative proteomics

To reveal the molecular targets of HPI-1 we designed a proteolysis-targeting chimeras (PROTACs) approach, that involves heterobifunctional molecules consisting of two-headed small ligands which bind to two different proteins; an E3 ligase and a target protein (POI) that is to be degraded through the ubiquitin-proteasome system³. The POI can eventually be identified by mass spectrometry-based proteomics. Using this strategy, we discovered the BET bromodomains as the targets of HPI-1, extending the scope of PROTACs as a novel target deconvolution technique.

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