# GalChimia

## Development of novel drug candidates for selective inhibition of Bcl-2 protein–protein interactions

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### Introduction

Protein-protein interactions as a therapeutic target

> The Bcl-2 family of proteins is central to the regulation of apoptosis, and their overexpression has been linked to certain types of lymphoma and carcinoma,<sup>1</sup> as well as resistance to conventional antitumor treatments. In particular, the interactions between pro- and antiapoptotic Bcl-2 family proteins control the integrity of the outer mitochondrial membrane.



#### Generation of Virtual Hits

HQL Pharmaceuticals

- > ChemSpaceScanner (CSS): Proprietary computational drug discovery platform for virtual screening of huge chemical libraries (>10<sup>30</sup> molecular entities) at high resolution.
- > A virtual library of potential candidates was designed and refined following ADMET and stability criteria. The structures were then clustered by scaffold similarity and further refined by synthetic feasibility and patentability.
- > The aim of this project was to develop a set of novel anti-cancer drug candidates that function by selectively inhibiting such protein–protein interactions (PPI).<sup>2</sup>
- > The present work focuses on the Hit Finding and Hit-to-lead phases for the development of novel inhibitors of PPI within the Bcl-2 protein family.



- > The obtained structures presented high chemical diversity, with very different scaffolds (bicyclic systems or highly functionalized aromatic rings) and varied substitution patterns including chiral centers in some cases.
- Activity optimization was performed through the typical designsynthesis-assay cycles with subsequent refinement of the computational models.



## Screening and Optimization

Synthetic routes were designed for each of the families, representative compounds were prepared, and their inhibitory activity evaluated by Surface Plasmon Resonance (SPR). Peptides corresponding to the BH3 domain helixes of pro-apoptotic Bcl-2 family members were attached to carboxyl-coated SPR chips via their *N*-terminus, and the binding of the protein to the chip-bound peptide was measured. Compounds were incubated at multiple concentrations with the protein prior to injection. \*Series HQL\_50 was added after the initial scaffold screening. (IC50 data shown only for the best compound within each series and iteration.)

| HQL_23   | HQL_3   | HQL_50*  | HQL_73  | R <sub>5</sub><br>HQL_87                                      | HQL_99                 | -1.00<br>-2.00<br>-3.00                                  |   |
|--|---|--|---|---|------------------------|--|---|
| Scc<br>scre                                    | affold<br>ening » »                               | <b>» » » » » » » » » » Ro</b> u                            | und 2 » » » » »   | » » » » » » Rou   | and 3 »»»»»            | »»»»»»»»»»   | ad  |
| 16 compounds                                   |   | 33 cor   | 33 compounds  |   | 62 compounds           |  |   |
| Series   | IC50 (µM)   | Series   | IC50 (µM)   | Series  | IC50 (µM)              | HQL_50   | D(123)  |
|  | not active  | HQL_32   | 2,6   | HQL_32  | 1,3                    | Bcl-xL   | 250 nM  |
| HQL_23   |   |  |   |   |                        | CYP1A2   | not active                                      |
| HQL_23<br>HQL_32                               | 32 – 200  | HQL 73   | 4 – 50  | HQL_73  | 1,5 – 12               |  |   |
| HQL_23<br>HQL_32                               | 32 – 200<br>180 - 300                             | HQL_73   | 4 - 50  | HQL_73  | 1,5 - 12<br>0.25 - 0.5 | CYP2C9   | 282 nM  |
| HQL_23<br>HQL_32<br>HQL_73                     | 32 – 200<br>180 – 300                             | HQL_73<br>HQL_50   | 4 – 50<br>0,5 – 2   | HQL_73<br>HQL_50  | 1,5 – 12<br>0,25 – 0,5 | CYP2C9<br>CYP2C19  | 282 nM<br>2,13 μM                               |
| HQL_23<br>HQL_32<br>HQL_73<br>HQL_87           | 32 – 200<br>180 – 300<br>not active               | HQL_73<br>HQL_50   | 4 - 50<br>0,5 - 2   | HQL_73<br>HQL_50  | 1,5 – 12<br>0,25 – 0,5 | CYP2C9<br>CYP2C19<br>Permeability (Papp)                 | 282 nM<br>2,13 μM<br>4,79 ×10 <sup>-6</sup> cm/ |
| HQL_23<br>HQL_32<br>HQL_73<br>HQL_87<br>HQL_99 | 32 – 200<br>180 – 300<br>not active<br>not active | HQL_73<br>HQL_50<br>New scaffold HQ<br>» » » » Stability i | 4 – 50<br>0,5 – 2<br>QL_50 very active.<br>ssues in HQL_32 solved b | HQL_73<br>HQL_50<br>by elongating linker in Ar <sub>2</sub> . | 1,5 – 12<br>0,25 – 0,5 | CYP2C9<br>CYP2C19<br>Permeability (Papp)<br>Cell survivo | 282 nM<br>2,13 μM<br>4,79 ×10 <sup>-6</sup> cm/ |



#### Conclusions

The aim of this project was to develop a set of novel anti-cancer drug candidates that function by selectively inhibiting Bcl-2 PPI. In collaboration with HQL Pharmaceuticals, a virtual library of potential candidates was designed. The virtual hits were clustered into families and selected compounds were chosen for synthesis. The activity of the compounds was evaluated by SPR. After the initial hit-finding stage, the structures were further optimized to improve the activity from micromolar to the nanomolar range. The selected lead was further evaluated in ADME/Tox assays.

Keywords: protein–protein interactions, Bcl-2, hit-to-lead, drug discovery



<sup>1</sup> Otake Y., Soundararajan S., Sengupta T.K., Kio E.A., Smith J.C., Pineda-Roman M., Stuart R.K., Spicer E.K., Fernandes D.J., *Blood*, 2007, 1:109, 3069.

<sup>2</sup> Mullard, A. Nat. Rev. Drug Discov., 2012, 11, 173.



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