Rapid Technique
For New Scaffold Generation
Tim Cheeseright, Martin J. Slater, Mark Mackey

Cresset BioMolecular Discovery Ltd, Biopark, Broadwater Road, Welwyn Garden City, Herts, AL7 3AX, UK  www.cresset-group.com

Introduction
Scaffold hopping remains a central task in medicinal chemistry for generating and protecting intellectual property. We present sparkV10, an application for rapidly generating reasonable yet novel scaffold replacements.

Our technique uses the molecular interaction fields of the parent molecule and assesses replacements in the context in which they will be synthesized. This enables the differing steric and electronic effects of potential new scaffolds to be used. An added bonus of our method is that replacements for terminal substituents can be considered alongside more central moieties enabling its use in growing fragments and lead optimization as well as lead generation.

We present the methodology (bottom) together with applications to core replacement (right) and to fragment growing/optimization (below).

Application to Fragment Growth
Search fragments from commercial compounds for moieties that create interactions with hinge region of p38, starting with a DFG-out binding region.

Search fragments from commercial compounds. Only consider replacements with < 3 rotatable bonds and attached through N or O, must contain Ar ring.

Experiment - grow 3K3I ligand to make interactions of 3ROC ligand - score replacements against both ligands (80% weight 3ROC, 20% weight 3K3I). Search fragments from commercial compounds. Only consider replacements with < 3 rotatable bonds and attached through N or O, must contain Ar ring.

Methodology
Our approach is encoded in a desktop software application “sparkV10”. Using sparkV10, a user identifies a region of a known active molecule that they wish to replace. The number of bonds broken by removing this piece is recorded together with the distance (δ₁ - δₙ) and angle (α₁ - αₙ) between any pair of broken bonds.

The angle and distance criteria are used, together with the number of connection points, to search a database of fragment conformations for replacement moieties. Matching fragment conformations are merged with the retained portions of the starter molecule to create a new “product” molecule. The product molecule is energy minimised and then scored as a replacement.

Scoring is performed using an average of field similarity and shape similarity on the product molecule. By default the scoring reflects the change relative to the original starter molecule but the user can choose to add other molecules that can be used in the scoring. In this way compounds with sub-optimal interactions can be improved by mimicking other known actives.

References