Statistical analysis of 3D distribution of ligand atoms in Protein Data Bank using kernel density estimation

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Scoring functions have been widely used to evaluate protein-ligand interaction in virtual screening. With increasing protein-ligand information in Protein Data Bank (PDB), various empirical and knowledge-based scoring functions are reported[1]. Previous studies of empirical scoring function, such as PMF score[2], used the structural information in PDB to train distance-dependent interaction energies between protein-ligand atom pairs. In this study, we analyzed the PDB data to estimate the 3D probability density distributions of ligand atoms for every grid points around amino acid residues, rather than to assess the interaction only considering the atom pair distance (Figure 1). Using kernel density estimation (KDE)[3], the probability distributions of ligand atoms around 20 amino acids were estimated for each of the atom types defined by PATTY[4]. As the result, the p-values were calculated for every grid points, and then compared to the score by force field calculation.


Figure 1. Brief procedure of the estimation of 3D distribution of ligand atoms.