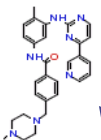
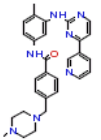
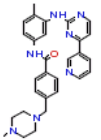


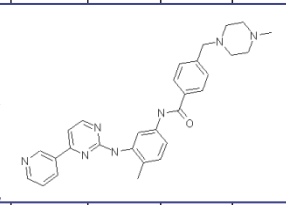
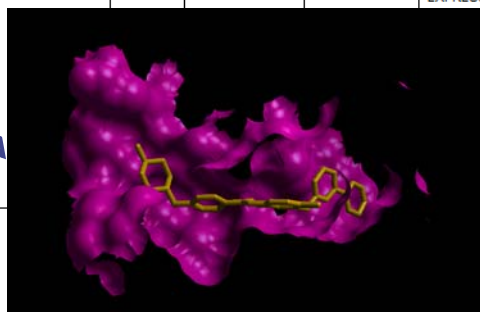
# From Ligand Query to Sites to New Ligand Ideas

The workflow consists of the following steps:

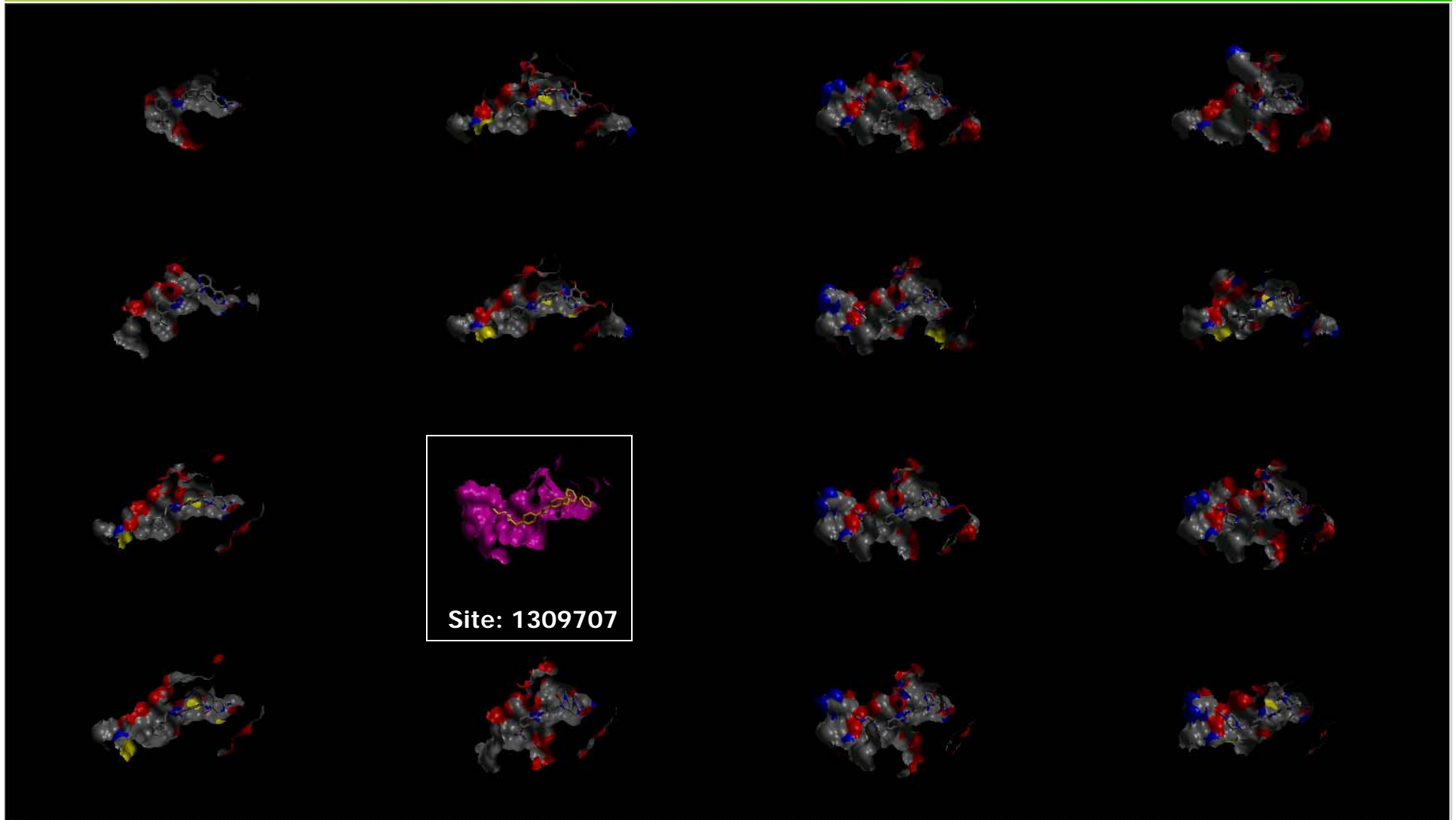
- Ligand Query:** A chemical structure of a benzamide derivative is shown with a distance of 9-11 Å between a nitrogen atom and an oxygen atom.
- Protein Structure:** A protein structure is shown with a specific site highlighted in pink.
- Protein-Ligand Complexes:** A grid of protein-ligand complexes is shown, illustrating the binding of various ligands to the protein site.
- Ligand Query Results:** Two windows displaying a grid of ligand structures, including:
  - STI\_4 (4-METHYLPIPERAZ...
  - BAX\_4 (4-[1][4]-CHLORO-3...
  - 460\_2 (5-(6-METHYLPIRID...
  - JRC\_6 (2,6-DICHLOROPHENY...
  - FRIC\_R ROSCOV...
  - AAZ\_4 (4-ARYL-2-PHENY...
  - STI\_BAX\_4, STI\_AAX\_7, STI\_BN\_7, BAX\_AAX\_3
  - BAX\_AAX\_1, STI\_BN\_5, STI\_BAX\_9, STI\_BAX\_7
  - STI\_BAX\_6, STI\_BAX\_2, STI\_BAX\_5, STI\_BAX\_9
  - STI\_4 (4-METHYLPIPERAZ...
  - STI\_BN\_2, STI\_AAX\_5, STI\_BN\_4

# Step 1: Find Co-complexes and Sites from Ligand-Structure-Search

Molecule	ligname	similarity	pdcode	sitoid	FourCode	pdid	pdBnxNumber	proteinid	title	classification	source	compound	releaseDate	journalTitle	journalReference	exctype
	STI	1	2pl0A	1309707	2pl0	2pl0	1305799	42526	LCK BOUND TO IMATINIB	TRANSFERASE	MOL_ID: 1; ORGANISM_SCIENTIFIC: HOMO SAPIENS; ORGANISM_COMMON: HUMAN; GENE: LCK; EXPRESSION_SYSTEM: SPODOPTERA FRUGIPERDA; EXPRESSION_SYSTEM_COMMON: FALL ARMYWORM; EXPRESSION_SYSTEM_VECTOR_TYPE: ... ON_SYSTEM_PLASMID:	MOL_ID: 1; MOLECULE: PROTO-ONCOGENE TYROSINE-PROTEIN KINASE LCK; CHAIN: A; FRAGMENT: PROTEIN KINASE; SYNONYM: P56-LCK, LYMPHOCYTE CELL-SPECIFIC PROTEIN-TYROSINE KINASE, LSK, T CELL-SPECIFIC PROTEIN-TYROSINE KINASE; EC: 2.7.10.2; ENGINEERED: YES	09-OCT-07	CLASSIFYING PROTEIN KINASE STRUCTURES GUIDES USE OF SELECTIVITY PROFILES TO PREDICT INACTIVE CONFORMATIONS: STRUCTURE OF LCK/IMATINIB COMPLEX	PROTEINS 2007	XRAY DIFFRACTION
	STI	1	2oiqA	1146914	2oiq	2oiq	1125109	26318	STRUCTURE OF CHICKEN C-SRC KINASE DOMAIN IN COMPLEX WITH THE CANCER DRUG IMATINIB.	TRANSFERASE	...; ORGANISM_SCIENTIFIC: GALLUS; ...M_COMMON: CHICKEN; GENE: SRC; EXPRESSION_SYSTEM: ESCHERICHIA COLI; EXPRESSION_SYSTEM_COMMON: BACTERIA; EXPRESSION_SYSTEM_STRAIN: BL21DE3; EXPRESSION_SYSTEM_VECTOR_TYPE: PLASMID; EXPRESSION_SYSTEM_PLASMID: PET28	MOL_ID: 1; MOLECULE: PROTO-ONCOGENE TYROSINE-PROTEIN KINASE SRC; CHAIN: A, B; FRAGMENT: KINASE DOMAIN; SYNONYM: P60-SRC, C-SRC, PP60C-SRC; EC: 2.7.10.2; ENGINEERED: YES	20-MAR-07	C-SRC BINDS TO THE CANCER DRUG IMATINIB WITH AN INACTIVE ABL/C-KIT CONFORMATION AND A DISTRIBUTED THERMODYNAMIC PENALTY.	STRUCTURE V. 15 299 2007	XRAY DIFFRACTION
	STI	1	2hyyA	918207	2hyy	2hyy	904013	16961	HUMAN ABL KINASE DOMAIN IN COMPLEX WITH IMATINIB (ST1571, GLIVEC)	TRANSFERASE	MOL_ID: 1; ORGANISM_SCIENTIFIC: HOMO SAPIENS; ORGANISM_COMMON: HUMAN; GENE: ABL1; EXPRESSION_SYSTEM: SPODOPTERA FRUGIPERDA; EXPRESSION_SYSTEM_COMMON: FALL ARMYWORM	MOL_ID: 1; MOLECULE: PROTO-ONCOGENE TYROSINE-PROTEIN KINASE ABL1; CHAIN: A, B, C, D; SYNONYM: P150, C-ABL, ABELSON MURINE LEUKEMIA VIRAL ONCOGENE HOMOLOG 1; EC: 2.7.10.2;	16-JAN-07	STRUCTURAL BIOLOGY CONTRIBUTIONS TO THE DISCOVERY OF DRUGS TO TREAT CHRONIC MYELOGENOUS LEUKAEMIA.	ACTA CRYSTALLOGR., SECT. D V. 63 80 2007	XRAY DIFFRACTION



## Step 2: Find Other Receptor Sites from Site-Similarity Search

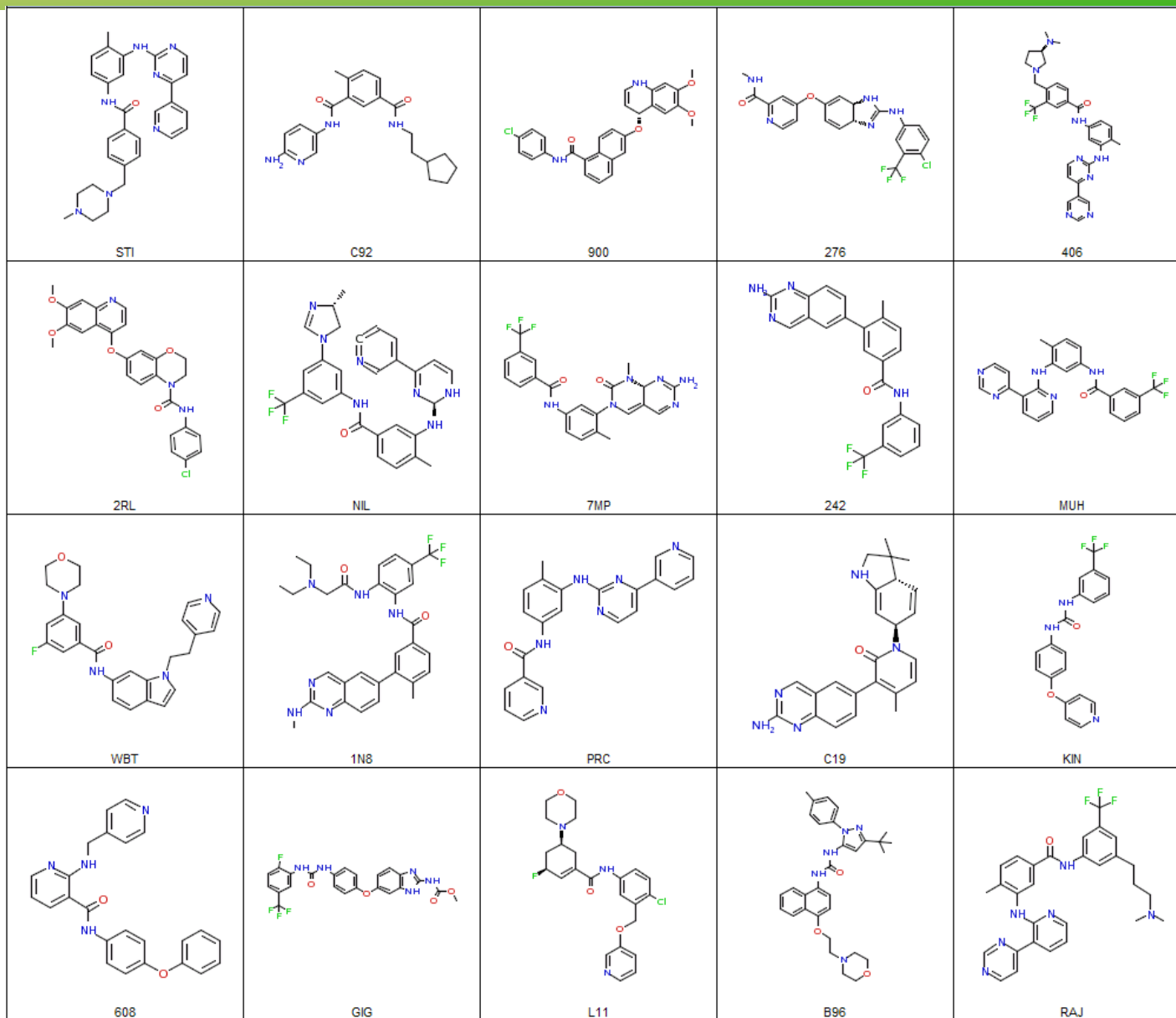


Chains	Chain Alignments	Sites	Site Alignments	
Site Name	Locus	Ligand	%Conf	Sequence Positions
pdb2pl0/s1309707 (chain A)	LCK	STI	100	.L.V.AVK.E.LM.L.LV.I.TEYM.GS.T.YIHR.L.IADF
pdb2of/s916548 (chain B)	LCK	242	100	.L.V.AVK.E.LM.L.LV.I.TEYM.G.S.I.V.H.L.IADF.I
pdb2rl5/s1396160 (chain A)	-	2RL	100	.LG.V.AVK.L.E.LL.I.VV.V.TEPCKPGL.L.CIB.LL.ICDF
pdb2e2b1/s1284639 (chain B)	ABL	406	100	.L.R.W.A.K.E.WM.H.LV.H.TEFMI.S.LL.FIHRD.LLVADF

## Example Site Similarity Results (Query: s1309707)

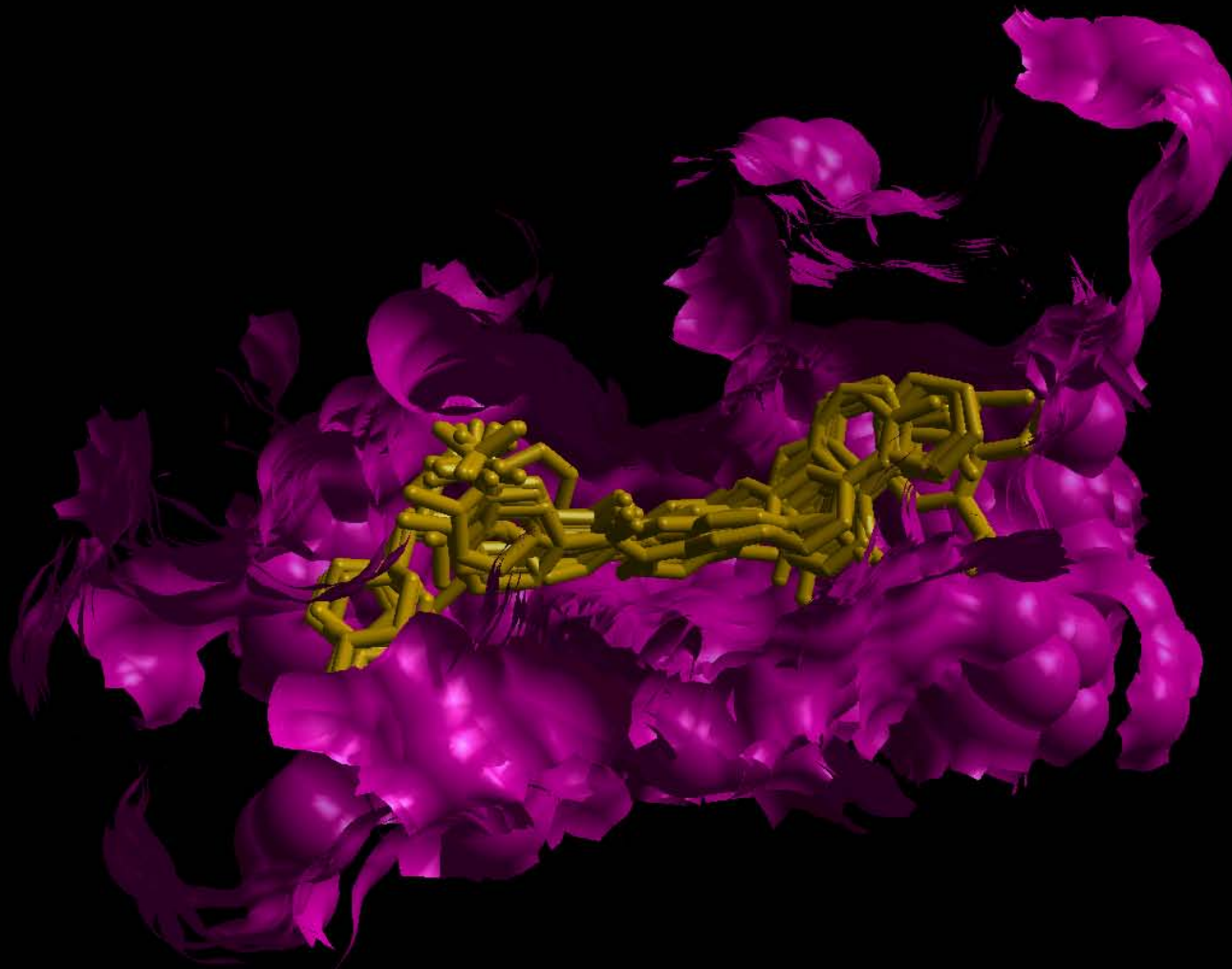
Site	SiteLigand	SiteProtein	SiteScore	ContactScore
1309707	STI	2pl0A	1000	1
1420904	C92	3cpbB	110.906	0.7
1384893	900	3b8qB	121.051	0.67
1322334	276	2qu5A	117.866	0.66
1284638	406	2e2bA	119.18	0.64
1396160	2RL	2rl5A	121.208	0.63
1400124	NIL	3cs9D	111.198	0.62
867405	7MP	2hiwA	101.948	0.61
916548	242	2ofvB	109.214	0.6
1147514	MUH	2oscA	104.115	0.6
776230	WBT	1wbtA	101.635	0.6
916805	1N8	2og8A	116.819	0.59
394066	PRC	1fpuB	107.297	0.57
1415780	C19	3cp9A	104.078	0.56
911671	KIN	2hznA	106.08	0.56
1148488	608	2p2iB	109.41	0.55
1300447	GIG	2oh4A	110.471	0.53
1320735	857	2qu6B	116.424	0.52
437653	B96	1kv2A	107.323	0.52
691631	L11	1w83A	101.268	0.52
1147212	RAJ	2oo8X	104.058	0.52
910098	GIN	2hz0B	108.713	0.51
1396708	P38	3bv2A	124.962	0.51
436174	BMU	1kv1A	88.568	0.5
1412158	G2G	2puuA	118.296	0.5
775147	L13	1wbvA	85.135	0.5
1415688	C52	3cpcB	102.25	0.48
1431710	GK6	3d83A	104.164	0.48

# Example Ligands Extracted from Similar Sites



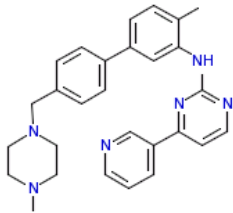
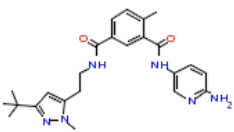
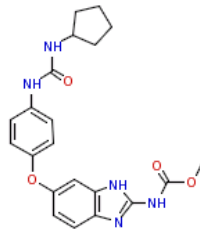
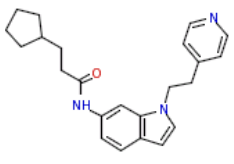
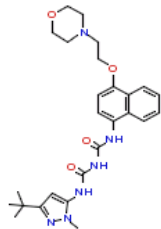
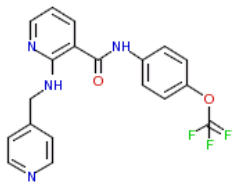
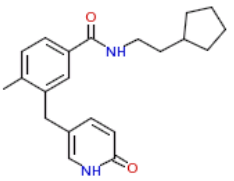
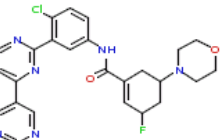
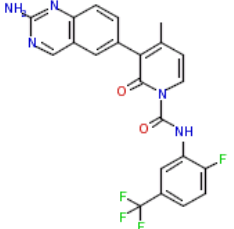
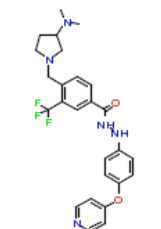
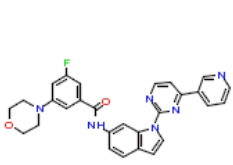
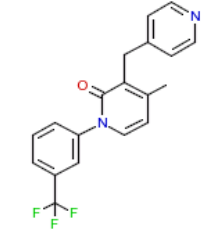
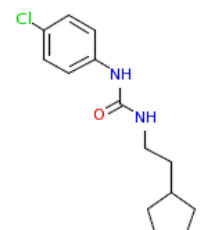
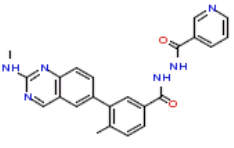
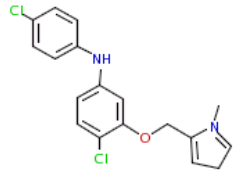
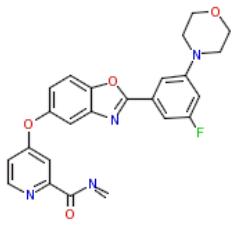
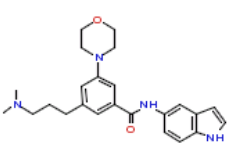
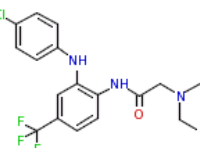
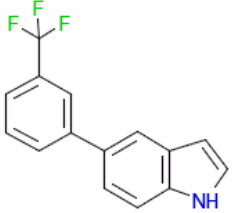
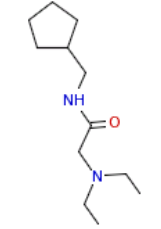


## Step 3: LigandCross – Mixing Ligand Features from Aligned Sites

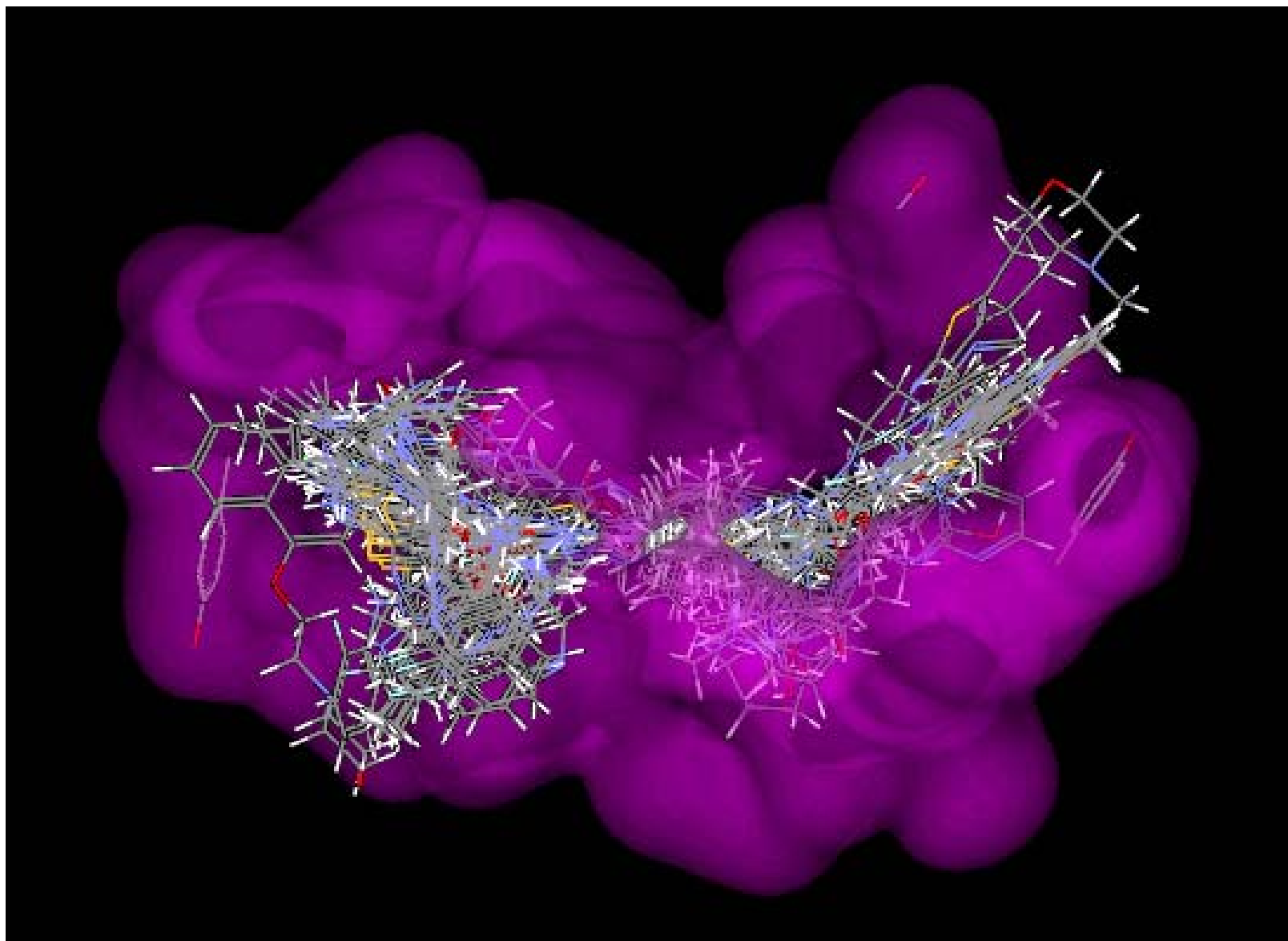


Chains	Chain Alignments	Sites	Site Alignments	
Site Name	Locus	Ligand	%Conf	Sequence Positions
pdb2pl0/s1309707 (chain A)	LCK	STI	100	.L.V.AVK.E.LM.D.LV.I.TEYM.GS.I.YIHR.L.IADF
pdb2of/s916548 (chain B)	LCK	242	100	.L.V.AVK.E.LM.D.LV.I.TEYM.G.I.V.H.L.IADF.I
pdb2rl5/s1396160 (chain A)	-	2RL	100	.LG.V.AVK.L.E.II.I.VV.V.TEFCKFGN.L.CIH.L.ICDF
pdb2e2b1/s1284639 (chain B)	ABL	406	100	.L.V.V.A.K.E.VM.I.LV.I.TEFMT.G.L.FIHRD.L.VADF

# Example LigandCross Results

 <p>STI_PRC_2 0.667</p>	 <p>C92_BMU_5 0.635</p>	 <p>C92_GIG_3 0.633</p>	 <p>C92_WBT_1 0.625</p>	 <p>B96_BMU_2 0.623</p>
 <p>608_276_3 0.608</p>	 <p>C92_GIN_7 0.608</p>	 <p>406_L11_6 0.577</p>	 <p>GIG_C52_1 0.574</p>	 <p>406_KIN_2 0.545</p>
 <p>NIL_WBT_6 0.538</p>	 <p>608_C52_2 0.529</p>	 <p>C92_BMU_1 0.520</p>	 <p>1N8_PRC_3 0.491</p>	 <p>857_BMU_4 0.480</p>
 <p>857_WBT_2 0.472</p>	 <p>RAJ_LI3_1 0.462</p>	 <p>1N8_BMU_2 0.449</p>	 <p>LI3_C52_2 0.385</p>	 <p>C92_1N8_1 0.375</p>

## Step 4: LigandCross Ligands reDocked into s1309707





# LigandCross Ligands with Reported Biological Activity

## Kinase Knowledgebase (pIC50)

## Bayesian Model Predictions (PP)

LC-ID	ABL	PDGFR	PDGFRB	JAK3	KDR	LCK	MAPK14	TEK	KIT	RAF1	ABL	PDGFR	PDGFRB	JAK3	KDR	LCK	MAPK14	TEK	KIT	RAF1
G2G_STI_12	6.7	8	8								0.40	0.90	0.76	0.81	0.59	0.15	0.89	0.45	0.70	0.37
900_STI_1	6.1	8	8								0.38	0.91	0.76	0.72	0.55	0.16	0.88	0.42	0.71	0.55
7MP_1N8_4				7.8	9	9.5	8.7				0.36	0.49	0.34	0.32	0.94	1.00	0.95	0.67	0.86	0.39
7MP_1N8_2				6.8	8.3	9.5	9				0.37	0.46	0.31	0.44	0.92	1.00	0.92	0.69	0.84	0.45
7MP_RAJ_3					8.4			8.4			0.35	0.73	0.50	0.49	0.92	0.81	0.86	0.94	0.74	0.37
7MP_GIN_4					7.6						0.16	0.50	0.40	0.82	0.95	0.67	0.70	0.41	0.76	0.51
242_C52_2									7.9		0.30	0.28	0.29	0.74	0.80	0.66	0.74	0.31	1.00	0.43
LI3_L11_1							7.2				0.31	0.73	0.55	0.84	0.74	0.69	0.62	0.36	0.76	0.85
608_GIG_7										6.1	0.28	0.61	0.57	0.69	0.93	0.50	0.60	0.68	0.85	0.50
KIN_BMU_4										6.1	0.31	0.43	0.45	0.78	0.75	0.57	0.77	0.33	0.81	0.25
G2G_KIN_3										6.1	0.25	0.51	0.52	0.75	0.89	0.59	0.64	0.43	0.84	0.43

