

## Supplemental Files

### **LASSO-ing Potential Nuclear Receptor Agonists and Antagonists: A New Computational Method For Database Screening**

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DISCLAIMER: This document has been subjected to review by the US Environmental Protection Agency and approved for publication.

**Running Head:** Models for Nuclear Receptor Agonists and Antagonists

**Supplemental Table 1.** LASSO 23 Surface Point Types with description.

Number	Surface Point Type	Description
1	Metal	positively charged metal ion interaction point
2	Charged_Hplus	positively charged hydrogen bond donor, e.g. Arginine
3	Primary_Amine_Hlp	primary amine hydrogen/lone-pair, e.g. $-\text{NH}_3^+$ or $-\text{NH}_2$
4	Hdonor	strong hydrogen bond donor H (polar-atom-H)
5	Weak_Hdonor	weak (secondary) hydrogen bond donor H (polarized C-H)
6	Charged_Lonepair	lone pair of negatively charged group, e.g. $\text{PO}_3$
7	Acid_Lonepair	lone pair of an acidic functional group, e.g. carboxolate
8	Lonepair strong	(primary) hydrogen bond acceptor lone pair
9	Weak_Lonepair	weak (secondary) hydrogen bond acceptor lone pair
10	Ambivalent_Hlp	donor H or acceptor Lp depending on protonation state
11	Rotatable_H	rotatable-hydroxy donor H
12	Rotatable_Lp	rotatable-hydroxy acceptor Lp
13	Hydrophob	H on $\text{Sp}^3$ hydrophobic carbon
14	H_Arom_Edge	H on hydrophobic carbon in aromatic ring (non-polarized)
15	Ws_Lipo	H on weak secondary hydrophobic atom (e.g. carbon next to polar)
16	Neutral	H/Lp on neutral atom (no recognized activity)
17	Pi_Aromatic	$\pi$ electron of an aromatic ring
18	Pi_Reson_Polar	$\pi$ electron on polar atom (N/O) in resonance chain, e.g. amide
19	Pi_Reson_Carbon	$\pi$ electron on carbon atom in resonance chain, e.g. amide
20	Pi_Sp2_Polar	$\pi$ electron on $\text{sp}^2$ polar atom (N/O)(non-resonating, nonaromatic)
21	Pi_Sp2_Carbon	$\pi$ electron on $\text{sp}^2$ carbon atom (non-resonating, non-aromatic)
22	Halogen	lone electron pair of a halogen atom (F,Cl,I,Br)
23	Sulfur	lone electron pair of a sulfur atom

## Supplemental Table 2

Model building details for PXR.

#Actives/ from position	1 <sup>st</sup> position	9 <sup>th</sup> position	17 <sup>th</sup> position	33 <sup>rd</sup> position	64 <sup>th</sup> position
8 actives	Model 8 from 1	Model 8 from 9	Model 8 from 17	Model 8 from 33	Model 8 from 64
16 actives	Model 16 from 1	Model 16 from 9	Model 16 from 17	Model 16 from 33	Model 16 from 64
32 actives	Model 32 from 1	Model 32 from 9	Model 32 from 17	Model 32 from 33	Model 32 from 64
64 actives	Model 64 from 1	Model 64 from 9	Model 64 from 17	Model 64 from 33	Model 64 from 64
128 actives	Model 128 from 1	Model 128 from 9	Model 128 from 17	Model 128 from 33	Model 128 from 64

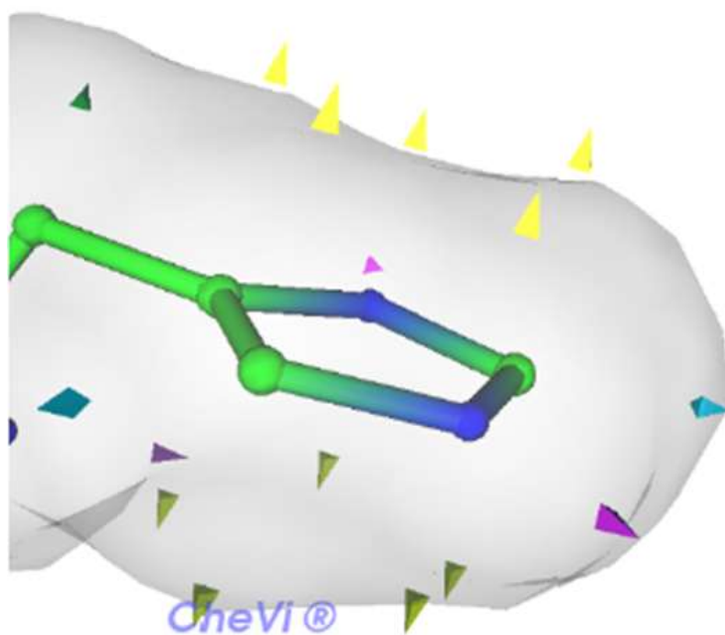
### Supplemental Table 3

#### LASSO 6.1 surface point types with Mibolerone's values:

<u>ID</u>	<u>Count</u>	<u>Name</u>	<u>Description</u>
1	0	Metal	positively charged metal ion point
2	0	Charged_Hplus	positively charged hydrogen, e.g. Argini
3	0	Primary_Amine_Hlp	primary amine hydrogen/lone-pair, e.g.
4	0	Hdonor	strong (primary) hydrogen bond donor t
5	0	Weak_Hdonor	weak (secondary) hydrogen bond dono
6	0	Charged_Lonepair	lone pair of negatively charged group, e
7	0	Acid_Lonepair	lone pair of an acid group, e.g. carboxo
<b>8</b>	<b>2</b>	<b>Lonepair strong</b>	<b>strong (primary) hydrogen bond acc</b>
9	0	Weak_Lonepair	weak (secondary) hydrogen bond acce
10	0	Ambivalent_Hlp	donor H OR acceptor Lp depending on
11	1	Rotatable_H	rotatable-hydroxy donor H
<b>12</b>	<b>2</b>	<b>Rotatable_Lp</b>	<b>rotatable-hydroxy acceptor Lp</b>
<b>13</b>	<b>28</b>	<b>Hydrophob</b>	<b>H on aliphatic (chain) hydrophobic c</b>
14	0	H_Arom_Edge	H on hydrophobic carbon in aromatic rii
15	0	Ws_Lipo	H on weak secondary hydrophobic aton
16	1	Neutral	H/Lp on neutral atom (no recognised ac
17	0	Pi_Aromatic	Pi electron of an aromatic ring
18	0	Pi_Reson_Polar	Pi electron on polar atom (N/O) in resor
19	1	Pi_Reson_Carbon	Pi electron on carbon atom in resonanc
20	1	Pi_Sp2_Polar	Pi electron on sp2 polar atom (N/O) (nc
<b>21</b>	<b>2</b>	<b>Pi_Sp2_Carbon</b>	<b>Pi electron on sp2 carbon atom (non</b>
22	0	Halogen	lone electron pair of a halogen atom (F,
23	0	Sulfur	lone electron pair of a sulfur atom
24	0	misc	

### Supplemental Figure 1

Histidine-like fragment of a molecules showing the surface point types as different colored triangles.



## Supplemental Figure 2

Mibolerone displayed in SimBioSys' CheVi 3D desktop visualization tool, showing the 3D structure, color-coded interaction surface of the molecule, and the surface points (triangles colored according to interaction types).

