High-Throughput Extraction of Interfacial Electrostatic Features from GLP-1-GLP-1R Complex Structures: A GLP-1-GLP-1R Based Mini GIBAC Perspective

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Abstract

From a structural point of view, understanding interfacial electrostatic features of ligand-receptor complexes is crucial for biophysics-based design of therapeutics with improved efficacy and safety. Here, with a high-throughput structural feature extraction approach, this article reports a set of electrostatic structural features from glucagon-like peptide-1 (GLP-1) and GLP-1 receptor (GLP-1R) complex structures determined using experimental tools. Leveraging a set of computational structural biophysical analyses, this article systematically characterized the electrostatic interactions within the GLP-1-GLP-1R complexes with two sets of criteria to identify interfacial salt bridges and hydrogen bonds. Overall, the results here reveal intricate details of electrostatic interactions critical for GLP-1 binding to its receptor GLP-1R, providing biophysical insights into the stabilization of the GLP-1-GLP-1R complex structure and the molecular basis of GLP-1/GLP-1R signaling. Furthermore, the electrostatic structural features extracted from GLP-1 and GLP-1R complex structures are useful in the development of a machine learning-based ligand-receptor binding affinity calculating model, i.e., a GLP-1-GLP-1R-specific general intermolecular binding affinity calculator (GIBAC), paving the way for accelerated discovery and structure-based design of drug candidates targeting the GLP-1/GLP-1R system with improved efficacy and safety.

Keywords: GLP-1, GLP-1R, High-Throughput Feature Extraction, Electrostatic Structural Features, General Intermolecular Binding Affinity Calculator (Gibac)

1. Introduction

Direct binding and interaction between ligands and receptors plays a fundamental role in various biological processes, including signal transduction, membrane protein trafficking, cell communication, and drug pharmacology [1–4]. Understanding the intricate details of these interactions, particularly the structural features at the binding interface of ligand and receptor, is crucial for unraveling the underlying mechanisms and designing effective therapeutics [5-7]. Among the myriad of ligand-receptor systems, the glucagon-like peptide-1 (GLP-1) and its cognate receptor, GLP-1 receptor (GLP-1R), constitute a pivotal axis in metabolic regulation and insulin secretion [8-10]. GLP-1, an incretin hormone secreted by enteroendocrine intestinal L-cells, exerts its physiological effects by binding to GLP-1R, a G-protein-coupled receptor predominantly expressed in pancreatic b-cells, among other tissues [11–13]. Activation of GLP-1R initiates a signaling cascade leading to enhanced insulin release, inhibition of glucagon secretion, and regulation of satiety, making it an attractive target

for the treatment of type 2 diabetes and obesity [14–16].

To date, a wide range of experimental structures have been determined for the GLP-1/GLP-1R complex, constituting the structural basis of GLP-1/GLP-1R interactions, particularly the structural electrostatic features at the binding interface [17–19]. Here, with a set of computational structural biophysical analyses for currently available experimental GLP-1/GLP-1R complex structures, this article systematically characterized the electrostatic interactions within the GLP-1-GLP-1R complexes with two sets of criteria to identify interfacial salt bridges and hydrogen bonds. By structurally elucidating the electrostatic landscape of GLP-1/GLP-1R interactions, this article aims to provide valuable insights into the molecular basis of ligand recognition and receptor activation, with structural and biophysical implications for rational design and development of therapeutics with improved efficacy and safety targeting the GLP-1/GLP-1R axis.

2. Motivation

Thanks to the continued development of experimental structural biology and the half-a-century old Protein Data Bank (PDB) [20,21], a comprehensive structural biophysical (CSB) analysis becomes possible [22–24] for specific ligand-receptor [25–27], antigen-antibody [28] or enzyme-substrate [29–31] complex structures deposited in PDB, expanding our understanding of the structural and biophysical basis of their interfacial structural stability, and facilitating the design of drug analogues with improved affinity to their interacting partners [6,32].

Semaglutide is a human GLP-1 analogue with 94% structural homology with native human GLP-1, with 3 important modifications: an amino acid substitution at position 8 that makes it less susceptible to degradation by dipeptidyl peptidase-4; lysine acylation of the peptide backbone, with a spacer and C-18 fatty di-acid chain at position 26 that provides strong, specific binding to albumin; and another amino acid substitution at position 34, which prevents C=18 fatty di-acid binding at the wrong site [33]. Interestingly, a semaglutide analogue was for the first time reported with a simple Val27-Arg28 exchange in its peptide backbone in 2021 [7]. Semaglutide is a glucagon-like peptide 1 analog used for the treatment of patients with type 2 diabetes mellitus. With 94% sequence similarity to human GLP-1, semaglutide is a glucagonlike peptide-1 receptor (GLP-1R) agonist, which binds directly to GLP-1R, causing various beneficial downstream effects that reduce blood glucose. Specifically, the amino acid sequence of GLP-1 is listed in italics in fasta format as below,

Semaglutide Native

Haegtftsdvssylegqaakefiawlvrgrg

The amino acid sequence of the semaglutide analogue with a simple Val27-Arg28 exchange in its peptide backbone is listed in italics in fasta format as below,

Semaglutide Mutant

Haegtftsdvssylegqaakefiawlrvgrg

Of further interest, a new pair of structure-stabilizing interfacial

salt bridges between Glu104 of chain A (GLP-1R) and Arg27 of chain B (semaglutide) are entirely due to the Val27-Arg28 exchange in the semaglutide peptide backbone, according to the binding affinity of semaglutide and GLP-1R calculated using Prodigy [34,35]. Nonetheless, the design of the semaglutide analogue with a simple Val27-Arg28 exchange was on a manual basis of close naked-eve inspection of the GLP-1-GLP-1R binding interface 4ZGM [19,36], and hand-picked from a set of analogue candidates of semaglutide to ensure improved ligand-receptor binding affinity [34,35] with minimum modification (i.e., a simple Val27-Arg28 exchange) to the backbone of semaglutide. Here, this article employs a high-throughput approach for extraction of electrostatic structural features from GLP-1-GLP-1R complexes towards the construction of a GLP-1-GLP-1R-based mini general intermolecular binding affinity calculator (GIBAC) with adequate accuracy and efficiency [2]. Overall, the motivation here stems from the pressing need within the drug discovery and design community for efficient methods to assess intermolecular binding affinities. particularly in the context of peptide-receptor interactions. By focusing on GLP-1-GLP-1R complexes, pivotal targets in diabetes and obesity [37,38], this article aims to showcase the utility of our miniaturized GIBAC approach in rapidly elucidating key electrostatic interactions critical for ligand-receptor binding for the purpose of discovery and design of therapeutics for diabetes and obesity with improved efficacy and safety.

3. Materials and Methods

As of February 25, 2024, a total of 44 experimental structures have been deposited in Protein Data Bank (PDB) [20] as listed in Table 1, according to a text query: *QUERY: Polymer Entity Description* = "Glucagon-like peptide 1 receptor" of the Protein Data Bank [20]. Among them, only two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM [19,36], respectively, providing an accurate structural basis of the GLP-1-GLP-1R interaction specificity for subsequent comprehensive structural biophysical (CSB) analysis of the two structural models (two yellow rows in Table 1).

PDB ID	Structure Title
8JIS	Cryo-EM structure of the GLP-1R/GCGR dual agonist peptide15-bound human GLP-1R-Gs complex
8JIP	Cryo-EM structure of the GLP-1R/GCGR dual agonist MEDI0382-bound human GLP-1R-Gs complex
8JIR	Cryo-EM structure of the GLP-1R/GCGR dual agonist SAR425899-bound human GLP-1R-Gs complex
7X8R	Cryo-EM structure of the Boc5-bound hGLP-1R-Gs complex
7X8S	Cryo-EM structure of the WB4-24-bound hGLP-1R-Gs complex
7S15	GLP-1 receptor bound with Pfizer small molecule agonist
7RG9	cryo-EM of human Glucagon-like peptide 1 receptor GLP-1R in apo form
7RGP	cryo-EM of human Glucagon-like peptide 1 receptor GLP-1R bound to tirzepatide
7VBH	Cryo-EM structure of the GIPR/GLP-1R/GCGR triagonist peptide 20-bound human GLP-1R-Gs complex
7VBI	Cryo-EM structure of the non-acylated tirzepatide (LY3298176)-bound human GLP-1R-Gs complex
7LLL	Exendin-4-bound Glucagon-Like Peptide-1 (GLP-1) Receptor in complex with Gs protein
7LLY	Oxyntomodulin-bound Glucagon-Like Peptide-1 (GLP-1) Receptor in complex with Gs protein
7S1M	Ex4-D-Ala bound to the glucagon-like peptide-1 receptor/g protein complex (conformer 1)
7S3I	Ex4-D-Ala bound to the glucagon-like peptide-1 receptor/g protein complex (conformer 2)
7RTB	Peptide-19 bound to the Glucagon-Like Peptide-1 Receptor (GLP-1R)
7DUR	Cryo-EM structure of the compound 2-bound human GLP-1 receptor-Gs complex
7EVM	Cryo-EM structure of the compound 2-bound human GLP-1 receptor-Gs complex
7KI0	Semaglutide-bound Glucagon-Like Peptide-1 (GLP-1) Receptor in Complex with Gs protein
7KI1	Taspoglutide-bound Glucagon-Like Peptide-1 (GLP-1) Receptor in Complex with Gs Protein
7DUQ	Cryo-EM structure of the compound 2 and GLP-1-bound human GLP-1 receptor-Gs complex
7E14	Compound2_GLP-1R_OWL833_Gs complex structure
7LCI	PF 06882961 bound to the glucagon-like peptide-1 receptor (GLP-1R):Gs complex
7LCJ	PF 06882961 bound to the glucagon-like peptide-1 receptor (GLP-1R):Gs complex
7LCK	PF 06882961 bound to the glucagon-like peptide-1 receptor (GLP-1R)
6XOX	cryo-EM of human GLP-1R bound to non-peptide agonist LY3502970
6X18	GLP-1 peptide hormone bound to Glucagon-Like peptide-1 (GLP-1) Receptor
6X19	Non peptide agonist CHU-128, bound to Glucagon-Like peptide-1 (GLP-1) Receptor
6X1A	Non peptide agonist PF-06882961, bound to Glucagon-Like peptide-1 (GLP-1) Receptor
7C2E	GLP-1R-Gs complex structure with a small molecule full agonist
6VCB	Cryo-EM structure of the Glucagon-like peptide-1 receptor in complex with G protein, GLP-1 peptide and a positive allosteric modulator
6ORV	Non-peptide agonist (TT-OAD2) bound to the Glucagon-Like peptide-1 (GLP-1) Receptor
5OTT	Extracellular domain of GLP-1 receptor in complex with exendin-4 variant Gly2Hcs/Thr5Hcs
5OTU	Extracellular domain of GLP-1 receptor in complex with GLP-1 variant Ala8Hcs/Thr11Hcs
5OTV	Extracellular domain of GLP-1 receptor in complex with GLP-1 variant Ala8Cyc/Thr11Hcs
5OTW	Extracellular domain of GLP-1 receptor in complex with GLP-1 variant Ala8Hcs/Thr11Cys
5OTX	Extracellular domain of GLP-1 receptor in complex with GLP-1 variant Ala8Cys/Thr11Cys
6GB1	Crystal structure of the GLP1 receptor ECD with Peptide 11
6B3J	3.3 angstrom phase-plate cryo-EM structure of a biased agonist-bound human GLP-1 receptor-Gs complex
5NX2	Crystal structure of thermostabilised full-length GLP-1R in complex with a truncated peptide agonist at 3.7 Å resolution
5E94	Antibody-bound Glucagon-like Peptide-1 receptor extracellular domain
4ZGM	Crystal structure of Semaglutide peptide backbone in complex with the GLP-1 receptor extracellular domain
3IOL	Crystal structure of Glucagon-Like Peptide-1 in complex with the extracellular domain of the Glucagon-Like Peptide-1 Receptor
3C59	Crystal structure of the ligand-bound glucagon-like peptide-1 receptor extracellular domain
3C5T	Crystal structure of the ligand-bound glucagon-like peptide-1 receptor extracellular domain

Table 1. Experimentally determined GLP-1R-related structures in the Protein Data Bank (PDB [20]) as of February 25, 2024 with a QUERY: Polymer Entity Description = "Glucagon-like peptide 1 receptor".

In this table, the two structural models representing the complex structures of ligand-bound GLP-1R are highlighted in two yellow rows, i.e., PDB IDs: 3IOL [17] and 4ZGM [36].

Of the two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM [19,36], two sets of amino acid sequences are listed in italics in fasta format as below,

>3IOL_1|Chain A|Glucagon-like peptide 1 receptor|Homo sapiens (9606) GSHMRPQGATVSLWETVQKWREYRRQCQRSLTEDPP-PATDLFCNRTFDEYACWPDGEPGSFV
NVSCPWYLPWASSVPQGHVYRFCTAEGLWLQKDNSSLP-WRDLSECEESKRGERSSPEEQLLFLY

>3IOL_2|Chain B|Glucagon|Homo sapiens (9606) HAEGTFTSDVSSYLEGOAAKEFIAWLVKGRG

>4ZGM_1|Chain A|Glucagon-like peptide 1 receptor|Homo sapiens (9606)

RPQGATVSLWETVQKWREYRRQCQRSLTEDPPPATDLFCN-RTFDEYACWPDGEPGSFVNVSC PWYLPWASSVPQGHVYRFC-TAEGLWLQKDNSSLPWRDLSECEESKRGERSSPEEQLLFLY >4ZGM_2|Chain B|Semaglutide peptide backbone; 8Aib,34R-GLP-1(7-37)-OH|Homo sapiens

(9606)

HAEGTFTSDVSSYLEGQAAKEFIAWLVRGRG

With the amino acid sequence alignment as shown in Figure 1, it is

quite clear that the sequences GLP-1 for 3IOL [17,18] and 4ZGM [19,36] is at position 33 or 34, i.e., an exchange of lysine and arginine, while the GLP-1R sequences for 3IOL [17,18] and 4ZGM

[19,36] are in complete alignment, except for the presence of one N-terminal four residue fragment (GSHM) in 3IOL [17,18] and the absence of it (GSHM) at N-terminal in 4ZGM [19,36].

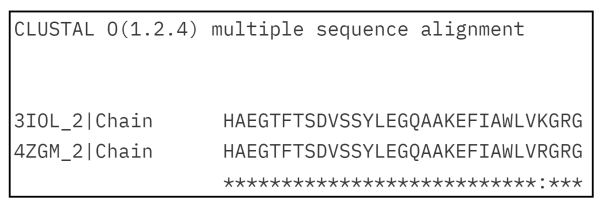


Figure 1: GLP-1 Amino Acid Sequence Alignment by Clustal Omega [39] Of Two Experimental Structures Represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM [19,36].

After the atomic coordinates file for PDB IDs: 3IOL [17,18] and 4ZGM [19,36] were downloaded from the PDB [20] website, Chimera [40] was employed to manually add hydrogen atoms to the structural model of the two structural models representing the GLP-1-GLP-1R complex structures, with file names 3IOH.pdb and 4ZGH.pdb, respectively. Afterwards, the two hydrogen-added structural models were subject to a set of comprehensive structural biophysical (CSB) analysis as described in [22] to identify key residue-specific interactions at the GLP-1-GLP-1R binding interface and uncover the interstructural biophysics underlying the GLP-1-GLP-1R complex structure. Specifically, the CSB analysis here [22] consists of the structural identification of salt bridges and side chain hydrogen bonds at the binding interface of GLP-1 and GLP-1R. Given the fact that native proteins are in dynamic equilibrium with their less-structured, partially folded and/or unfolded states [41], this article uses two sets of screening criteria for the structural identification of potential hotspots at the GLP-1-GLP-1R binding interface in the two structural models i.e., PDB IDs: 3IOL [17,18] and 4ZGM [19,36].

First, the same set of criteria (referred to as the old criteria below) as in [22] was used, i.e., the interfacial salt bridge analysis was conducted with an in-house python script only for titrateable residues (Asp, Glu, Lys, Arg and His), 4.0 Å was used as the cutoff distance for the two oppositely charged groups [22,42]. The hydrogen bond analysis was also conducted for only side chain nuclei with an in-house python script, and employed two geometric criteria: (a) a cutoff value of the angle formed by acceptor (A), donor (D) and hydrogen (H) ($\angle ADH$) of 30°; (b) a cutoff value of donor-acceptor distance at 3.0 Å. That is, a hydrogen bond is only considered to be formed if $\angle ADH$ is not larger than 30° and the donor-acceptor distance is not larger than 3.0 Å [22,42].

Afterwards, a new set of criteria (referred to as the new criteria

below) was used to account for the GLP-1-GLP-1R complex structures, i.e., the interfacial salt bridge analysis was conducted with an in-house python script only for titrateable residues (Asp, Glu, Lys, Arg and His), 6.0 Å was used as the cutoff distance for the two oppositely charged groups [22,42]. The hydrogen bond analysis was also conducted for only side chain nuclei with an inhouse python script, and employed two geometric criteria: (a) a cutoff value of the angle formed by acceptor (A), donor (D) and hydrogen (H) ($\angle ADH$) of 50°; (b) a cutoff value of donor-acceptor distance at 5.0 Å. That is, a hydrogen bond is only considered to be formed if $\angle ADH$ is not larger than 50° and the donor-acceptor distance is not larger than 5.0 Å [22,42].

Here, the in-house python scripts essentially are the same as those used in [43], except for the differences in three key parameters related to the screening criteria, i.e., the salt bridge distance cutoff in Å, cutoff angle $\angle ADH$ in ° for hydrogen bonding network screening, and the cutoff distance (in Å) of donor-acceptor for hydrogen bonding network screening.

4. Results

4.1. Residue-specific electrostatic interactions at the GLP-1-GLP-1R binding interface: a structural screening with the old criteria

For the two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM [19,36], a set of residue-specific electrostatic interactions at the GLP-1-GLP-1R binding interface were identified with a structural screening with the old criteria, including salt bridges at the GLP-1-GLP-1R binding interface for both 3IOL [17,18] and 4ZGM [19,36] as included in Table 2 and side chain hydrogen bonds at the GLP-1-GLP-1R binding interface for both 3IOL [17,18] and 4ZGM [19,36] as included in Table 3.

PDB ID	Residue A	Atom A	Residue B	Atom B	Distance (Å)
3IOH	A_ARG_44	NH2	A_GLU_41	OE1	3.682
3IOH	A_ARG_44	NH2	A_GLU_41	OE2	2.811
3IOH	A_ARG_64	NH2	A_ASP_74	OD1	3.801
3IOH	A_ARG_64	NH2	A_ASP_74	OD2	2.782
3IOH	A_ARG_102	NH1	A_ASP_67	OD2	3.054
3IOH	A_ARG_121	NH1	A_ASP_67	OD1	3.519
3IOH	A_ARG_121	NH1	A_ASP_67	OD2	2.959
3IOH	B_LYS_26	NZ	A_GLU_128	OE1	3.212
3IOH	B_LYS_26	NZ	A_GLU_128	OE2	2.924
4ZGH	A_LYS_38	NZ	A_GLU_34	OE1	3.823
4ZGH	A_ARG_44	NH2	A_GLU_41	OE1	3.861
4ZGH	A_ARG_44	NH2	A_GLU_41	OE2	2.747
4ZGH	A_ARG_64	NH2	A_ASP_74	OD1	3.779
4ZGH	A_ARG_64	NH2	A_ASP_74	OD2	2.832
4ZGH	A_ARG_102	NH1	A_ASP_67	OD2	3.198
4ZGH	A_ARG_121	NH1	A_ASP_67	OD1	3.561
4ZGH	A_ARG_121	NH1	A_ASP_67	OD2	2.953
4ZGH	B_LYS_26	NZ	A_GLU_128	OE1	3.409
4ZGH	B_LYS_26	NZ	A_GLU_128	OE2	2.771

Table 2: Salt bridging screening for two GLP-1-GLP-1R complex structures (PDB IDs: 3IOL [17,18] and 4ZGM [19,36]). In this table, the residue naming scheme is Chain ID_residue name_residue number.

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
3IOH	OE1, A_GLU_41	NE, A_ARG_44	HE, A_ARG_44	2.87	1.87	4.40
3IOH	OE2, A_GLU_41	NH2, A_ARG_44	HH21, A_ARG_44	2.81	1.86	15.77
3IOH	OE2, A_GLU_41	NE2, A_GLN_45	HE21, A_GLN_45	2.80	1.81	7.74
3IOH	OD2, A_ASP_74	NH2, A_ARG_64	HH22, A_ARG_64	2.78	1.78	5.75
3IOH	OD2, A_ASP_67	NE1, A_TRP_72	HE1, A_TRP_72	2.69	1.83	24.82
3IOH	OE2, A_GLU_125	OH, A_TYR_101	HH, A_TYR_101	2.63	1.69	8.92
3IOH	OD2, A_ASP_67	NH1, A_ARG_121	HH12, A_ARG_121	2.96	2.09	25.64
3IOH	OD1, A_ASP_122	OG, A_SER_124	HG, A_SER_124	2.65	1.73	12.69
4ZGH	OE1, A_GLU_41	NE, A_ARG_44	HE, A_ARG_44	2.92	2.08	10.96
4ZGH	OE2, A_GLU_41	NH2, A_ARG_44	HH21, A_ARG_44	2.75	1.91	11.18
4ZGH	OE2, A_GLU_41	NE2, A_GLN_45	HE21, A_GLN_45	2.99	2.14	5.73
4ZGH	OD1, A_ASP_74	NE, A_ARG_64	HE, A_ARG_64	2.97	2.12	3.34
4ZGH	OD2, A_ASP_74	NH2, A_ARG_64	HH21, A_ARG_64	2.83	1.98	7.04
4ZGH	OD2, A_ASP_67	NE1, A_TRP_72	HE1, A_TRP_72	2.80	2.02	20.99
4ZGH	OE2, A_GLU_125	OH, A_TYR_101	HH, A_TYR_101	2.82	1.99	6.37
4ZGH	OD2, A_ASP_67	NH1, A_ARG_121	HH11, A_ARG_121	2.95	2.17	20.60
4ZGH	OD1, A_ASP_122	OG, A_SER_124	HG, A_SER_124	2.67	1.91	21.21
4ZGH	OE2, A_GLU_128	NZ, B_LYS_26	HZ2, B_LYS_26	2.77	2.03	27.77
4ZGH	OXT, B_GLY_37	NE1, B_TRP_31	HE1, B_TRP_31	2.92	2.09	12.63

Table 3: Side chain hydrogen bonding network analysis for two GLP-1-GLP-1R complex structures (PDB IDs: 3IOL [17,18] and 4ZGM [19,36]). In this table, the residue naming scheme is Chain ID_residue name residue number, \angle ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (\angle ADH).

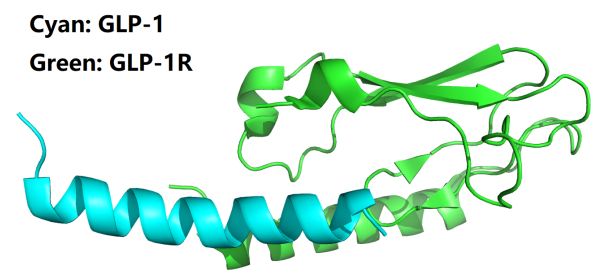


Figure 2: Structural model of the B27Arg-B28Val mutant of semaglutide peptide backbone (cyan cartoon) in complex with the GLP-1 receptor extracellular domain (green cartoon). This figure is prepared by PyMol [44] with PDB ID 3IOL [17,18].

To make it as clear and concise as possible, all interfacial salt bridges are tabulated in Table 4 for two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM [19,36]. Coincidentally, the two sets of interfacial salt bridges which contribute towards the stabilization of the complex structure of GLP-1 and GLP-1R are identical for

3IOL [17,18] and 4ZGM [19,36], i.e., two interfacial salt bridges (3.212 and 2.924 Å) between Lys26 of GLP-1 and Glu128 of GLP-1R for 3IOL [17,18], as shown by yellow dotted lines in Figures 3 and 4, and two interfacial salt bridges (3.409 and 2.771 Å) between Lys26 of semaglutide and Glu128 of GLP-1R for 4ZGM [19,36], as shown by yellow dotted lines in Figures 5 and 6.

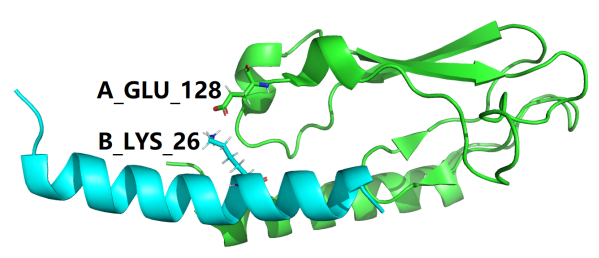


Figure 3: One interfacial salt bridge (Table 4) between semaglutide peptide backbone (cyan cartoon) in complex with the GLP-1 receptor extracellular domain (green cartoon). This figure is prepared by PyMol [44] with PDB ID 3IOL [17,18]. In this figure, the color scheme is the same as in Figure 2.

PDB ID	Residue A	Atom A	Residue B	Atom B	Distance (Å)
3IOH	B_LYS_26	NZ	A_GLU_128	OE1	3.212
3IOH	B_LYS_26	NZ	A_GLU_128	OE2	2.924
4ZGH	B_LYS_26	NZ	A_GLU_128	OE1	3.409
4ZGH	B_LYS_26	NZ	A_GLU_128	OE2	2.771

Table 4: Interfacial Salt bridging screening for two GLP-1-GLP-1R complex structures (PDB IDs: 3IOL [17,18] and 4ZGM [19,36]). In this table, the residue naming scheme is Chain ID_residue name_residue number.

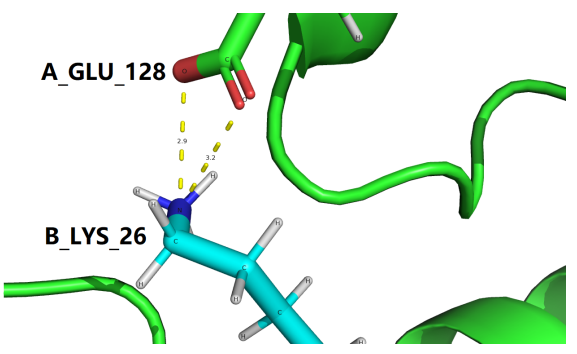


Figure 4: One interfacial salt bridge (yellow dotted lines, Table 4) between semaglutide peptide backbone (cyan cartoon) in complex with the GLP-1 receptor extracellular domain (green cartoon). This figure is prepared by PyMol [44] with PDB ID 3IOL [17,18]. In this figure, the color scheme is the same as in Figure 2.

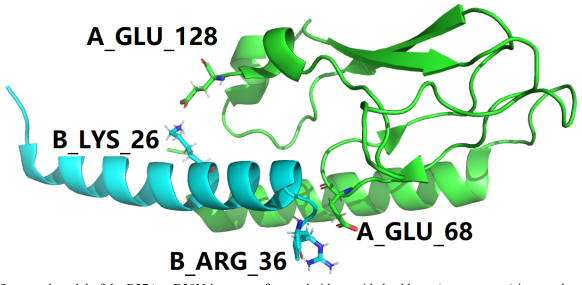


Figure 5: Structural model of the B27Arg-B28Val mutant of semaglutide peptide backbone (cyan cartoon) in complex with the GLP-1 receptor extracellular domain (green cartoon). This figure is prepared by PyMol [44] with PDB ID 4ZGM [19,36]. In this figure, the color scheme is the same as in Figure 2.

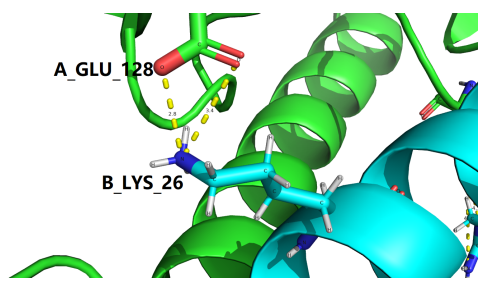


Figure 6: Structural model of the B27Arg-B28Val mutant of semaglutide peptide backbone (cyan cartoon) in complex with the GLP-1 receptor extracellular domain (green cartoon). This figure is prepared by PyMol [44] with PDB ID 4ZGM [19,36]. In this figure, the color scheme is the same as in Figure 2.

In addition to the structure-stabilizing interfacial salt bridges mentioned above, a set of side chain hydrogen bonds at the GLP-1-GLP-1R binding interface for both 3IOL [17,18] and 4ZGM [19,36] are also included in Tables 3. Seperately, for PDB ID 3IOL

[17,18], the set of side chain hydrogen bonds at the GLP-1-GLP-1R binding interface were included in Table 5, while for PDB ID 4ZGM [19,36], the set of side chain hydrogen bonds at the GLP-1-GLP-1R binding interface were included in Table 6.

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
3IOH	OE1, A_GLU_41	NE, A_ARG_44	HE, A_ARG_44	2.87	1.87	4.40
3IOH	OE2, A_GLU_41	NH2, A_ARG_44	HH21, A_ARG_44	2.81	1.86	15.77
3IOH	OE2, A_GLU_41	NE2, A_GLN_45	HE21, A_GLN_45	2.80	1.81	7.74
3IOH	OD2, A_ASP_74	NH2, A_ARG_64	HH22, A_ARG_64	2.78	1.78	5.75
3IOH	OD2, A_ASP_67	NE1, A_TRP_72	HE1, A_TRP_72	2.69	1.83	24.82
3IOH	OE2, A_GLU_125	OH, A_TYR_101	HH, A_TYR_101	2.63	1.69	8.92
3IOH	OD2, A_ASP_67	NH1, A_ARG_121	HH12, A_ARG_121	2.96	2.09	25.64
3IOH	OD1, A_ASP_122	OG, A_SER_124	HG, A_SER_124	2.65	1.73	12.69

Table 5: 3IOH-Specific Side Chain Hydrogen Bonding Analysis. In this table, the Residue Naming Scheme is Chain ID Residue name Residue Number, \angle ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (\angle ADH).

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
4ZGH	OE1, A_GLU_41	NE, A_ARG_44	HE, A_ARG_44	2.92	2.08	10.96
4ZGH	OE2, A_GLU_41	NH2, A_ARG_44	HH21, A_ARG_44	2.75	1.91	11.18
4ZGH	OE2, A_GLU_41	NE2, A_GLN_45	HE21, A_GLN_45	2.99	2.14	5.73
4ZGH	OD1, A_ASP_74	NE, A_ARG_64	HE, A_ARG_64	2.97	2.12	3.34
4ZGH	OD2, A_ASP_74	NH2, A_ARG_64	HH21, A_ARG_64	2.83	1.98	7.04
4ZGH	OD2, A_ASP_67	NE1, A_TRP_72	HE1, A_TRP_72	2.80	2.02	20.99
4ZGH	OE2, A_GLU_125	OH, A_TYR_101	HH, A_TYR_101	2.82	1.99	6.37
4ZGH	OD2, A_ASP_67	NH1, A_ARG_121	HH11, A_ARG_121	2.95	2.17	20.60
4ZGH	OD1, A_ASP_122	OG, A_SER_124	HG, A_SER_124	2.67	1.91	21.21
4ZGH	OE2, A_GLU_128	NZ, B_LYS_26	HZ2, B_LYS_26	2.77	2.03	27.77
4ZGH	OXT, B_GLY_37	NE1, B_TRP_31	HE1, B_TRP_31	2.92	2.09	12.63

Table 6: 4ZGH-specific side chain hydrogen bonding analysis. In this table, the residue naming scheme is Chain ID_residue name residue number, \angle ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (\angle ADH).

Among the two sets of side chain hydrogen bonds at the GLP-1-GLP-1R binding interface (Tables 5 and 6), one extraordinary interfacial side chain hydrogen bond was found to exist between the oppositely charged side chains of Lys26 of semaglutide and Glu128 of GLP-1R for 4ZGM [19,36], as described with the details in Table 7. Of further structural and biophysical interest, this outstanding interfacial side chain hydrogen bond is formed

between the same pair of oppositely charged amino acid residues as the pair of residues in two interfacial salt bridges (3.212 and 2.924 Å) between Lys26 of GLP-1 and Glu128 of GLP-1R for 3IOL [17,18], as shown by yellow dotted lines in Figures 3 and 4, and two interfacial salt bridges (3.409 and 2.771 Å) between Lys26 of semaglutide and Glu128 of GLP-1R for 4ZGM [19,36], as shown by yellow dotted lines in Figures 5 and 6.

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
4ZGH	OE2, A_GLU_128	NZ, B_LYS_26	HZ2, B_LYS_26	2.77	2.03	27.77

Table 7: 4ZGH-specific interfacial side chain hydrogen bonding analysis., the residue naming scheme is Chain ID_residue name residue number, \angle ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (\angle ADH).

4.2. Residue-specific electrostatic interactions at the GLP-1-GLP-1R binding interface: a structural screening with the new criteria

As discussed above in the section of Materials and Methods, native proteins are in dynamic equilibrium with their less-structured, partially folded and/or unfolded states [41], this article uses two sets of screening criteria for the structural identification of potential hotspots at the GLP-1-GLP-1R binding interface in the two structural models i.e., PDB IDs: 3IOL [17,18] and 4ZGM

[19,36]. Therefore, for the two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM, a set of residue-specific electrostatic interactions at the GLP-1-GLP-1R binding interface were identified with a structural screening with the new criteria, including salt bridges at the GLP-1-GLP-1R binding interface for both 3IOL and 4ZGM as included in Tables 8 and 9 and side chain hydrogen bonds at the GLP-1-GLP-1R binding interface for both 3IOL and 4ZGM [19,36] as included in Tables 10, 11, 12 and 13, and also shown in Figure 7.

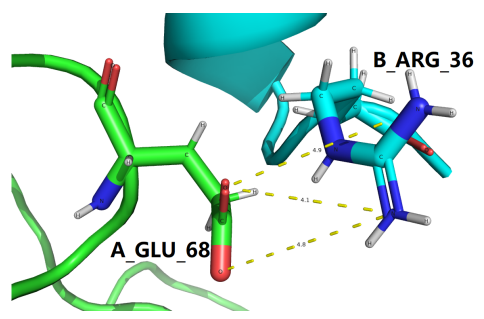


Figure 7: Structural model of the B27Arg-B28Val mutant of semaglutide peptide backbone (cyan cartoon) in complex with the GLP-1 receptor extracellular domain (green cartoon). This figure is prepared by PyMol [44] with PDB ID 4ZGM [19,36]. In this figure, the color scheme is the same as in Figure 2.

For the two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL and 4ZGM, a set of residue-specific electrostatic interactions at the GLP-1-GLP-1R binding interface were identified with a structural screening with the new criteria,

including salt bridges at the GLP-1-GLP-1R binding interface for both 3IOL [17,18] and 4ZGM [19,36] as included in Table 8 and side chain hydrogen bonds at the GLP-1-GLP-1R binding interface for both 3IOL and 4ZGM as included in Tables 10 and 11.

PDB ID	Residue A	Atom A	Residue B	Atom B	Distance (Å)
3IOH	A_LYS_38	NZ	A_GLU_34	OE1	5.019
3IOH	A_LYS_38	NZ	A_GLU_34	OE2	5.563
3IOH	A_ARG_43	NH1	A_GLU_68	OE1	5.108
3IOH	A_ARG_44	NH1	A_GLU_41	OE1	5.031
3IOH	A_ARG_44	NH1	A_GLU_41	OE2	4.860
3IOH	A_ARG_44	NH2	A_GLU_41	OE1	3.682
3IOH	A_ARG_44	NH2	A_GLU_41	OE2	2.811
3IOH	A_ARG_48	NH1	A_GLU_41	OE2	5.664
3IOH	A_ARG_64	NH1	A_ASP_53	OD1	5.728
3IOH	A_ARG_64	NH1	A_ASP_53	OD2	4.864
3IOH	A_ARG_64	NH1	A_ASP_74	OD1	5.246
3IOH	A_ARG_64	NH1	A_ASP_74	OD2	4.902
3IOH	A_ARG_64	NH2	A_ASP_53	OD2	5.347
3IOH	A_ARG_64	NH2	A_ASP_74	OD1	3.801
3IOH	A_ARG_64	NH2	A_ASP_74	OD2	2.782
3IOH	A_HIS_99	ND1	A_GLU_125	OE1	4.292
3IOH	A_HIS_99	ND1	A_GLU_125	OE2	5.603
3IOH	A_ARG_102	NH1	A_ASP_67	OD1	4.865
3IOH	A_ARG_102	NH1	A_ASP_67	OD2	3.054
3IOH	A_ARG_102	NH2	A_ASP_67	OD2	5.183
3IOH	A_ARG_121	NH1	A_ASP_67	OD1	3.519
3IOH	A_ARG_121	NH1	A_ASP_67	OD2	2.959
3IOH	A_ARG_121	NH2	A_ASP_67	OD1	5.721
3IOH	A_ARG_121	NH2	A_ASP_67	OD2	5.061
3IOH	B_LYS_26	NZ	A_GLU_128	OE1	3.212
3IOH	B_LYS_26	NZ	A_GLU_128	OE2	2.924
4ZGH	A_LYS_38	NZ	A_GLU_34	OE1	3.823
4ZGH	A_LYS_38	NZ	A_GLU_34	OE2	5.820
4ZGH	A_ARG_43	NH1	A_GLU_68	OE1	5.144
4ZGH	A_ARG_44	NH1	A_GLU_41	OE1	5.098
4ZGH	A_ARG_44	NH1	A_GLU_41	OE2	4.740
4ZGH	A_ARG_44	NH2	A_GLU_41	OE1	3.861
4ZGH	A_ARG_44	NH2	A_GLU_41	OE2	2.747
4ZGH	A_ARG_48	NH1	A_GLU_41	OE2	5.698
4ZGH	A_ARG_64	NH1	A_ASP_53	OD1	5.693
4ZGH	A_ARG_64	NH1	A_ASP_53	OD2	4.947
4ZGH	A_ARG_64	NH1	A_ASP_74	OD1	5.111
4ZGH	A_ARG_64	NH1	A_ASP_74	OD2	4.906
4ZGH	A_ARG_64	NH2	A_ASP_53	OD2	5.424
4ZGH	A_ARG_64	NH2	A_ASP_74	OD1	3.779
4ZGH	A_ARG_64	NH2	A_ASP_74	OD2	2.832
4ZGH	A_HIS_99	ND1	A_GLU_125	OE1	4.316
4ZGH	A_HIS_99	ND1	A_GLU_125	OE2	5.690
4ZGH	A_ARG_102	NH1	A_ASP_67	OD1	4.996
4ZGH	A_ARG_102	NH1	A_ASP_67	OD2	3.198
4ZGH	A_ARG_102	NH2	A_ASP_67	OD2	5.343
4ZGH	A_ARG_121	NH1	A_ASP_67	OD1	3.561
4ZGH	A_ARG_121	NH1	A_ASP_67	OD2	2.953
4ZGH	A_ARG_121	NH2	A_ASP_67	OD1	5.760
4ZGH	A_ARG_121	NH2	A_ASP_67	OD2	5.107

4ZGH	B_LYS_26	NZ	A_GLU_128	OE1	3.409
4ZGH	B_LYS_26	NZ	A_GLU_128	OE2	2.771
4ZGH	B_ARG_34	NH1	B_GLU_27	OE1	5.552
4ZGH	B_ARG_34	NH2	B_GLU_27	OE1	5.072
4ZGH	B_ARG_36	NH1	A_GLU_68	OE1	4.147
4ZGH	B_ARG_36	NH1	A_GLU_68	OE2	4.835
4ZGH	B_ARG_36	NH2	A_GLU_68	OE1	4.893

Table 8: Salt bridging screening with the new criteria as defined in the section of Materials and Methods for two GLP-1-GLP-1R complex structures (PDB IDs: 3IOL [17,18] and 4ZGM [19,36]). In this table, the residue naming scheme is Chain ID residue name residue number.

To make it as clear and concise as possible, all interfacial salt bridges are tabulated in Table 9 for two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM [19,36]. With a close comparison between Table 4 and Table 9, all four interfacial salt bridges in Table 4 are also present in Table 9. However, three additional interfacial salt bridges (according to the new criteria) are identified to be formed

between B ARG 36 of semaglutide backbone and A GLU 68 of GLP-1R in 4ZGM [19,36], as shown in Figures 5 and 7. Therefore, at the binding interface of semaglutide backbone and GLP-1R, two interfacial salt bridges are like two electrostatic clips [22], stabilizing the complex structure of semaglutide backbone and GLP-1R and facilitating the downstream signal related to the metabolism [33,38,45].

PDB ID	Residue A	Atom A	Residue B	Atom B	Distance (Å)
3IOH	B_LYS_26	NZ	A_GLU_128	OE1	3.212
3IOH	B_LYS_26	NZ	A_GLU_128	OE2	2.924
4ZGH	B_LYS_26	NZ	A_GLU_128	OE1	3.409
4ZGH	B_LYS_26	NZ	A_GLU_128	OE2	2.771
4ZGH	B_ARG_36	NH1	A_GLU_68	OE1	4.147
4ZGH	B_ARG_36	NH1	A_GLU_68	OE2	4.835
4ZGH	B_ARG_36	NH2	A_GLU_68	OE1	4.893

Table 9: Interfacial salt bridging screening with the new criteria as defined in the section of Materials and Methods for two GLP-1-GLP-1R complex structures (PDB IDs: 3IOL [17,18] and 4ZGM [19,36]). In this table, the residue naming scheme is Chain ID residue name residue number

In addition to the structure-stabilizing interfacial salt bridges 1-GLP-1R binding interface for both 3IOL [17,18] and 4ZGM mentioned above, a set of side chain hydrogen bonds at the GLP-

[19,36] are also included in Tables 10 and 11, respectively.

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
3IOH	OE1, A_GLU_41	NE, A_ARG_44	HE, A_ARG_44	2.87	1.87	4.40
3IOH	OE2, A_GLU_41	NH2, A_ARG_44	HH21, A_ARG_44	2.81	1.86	15.77
3IOH	OE2, A_GLU_41	NE2, A_GLN_45	HE21, A_GLN_45	2.80	1.81	7.74
3IOH	OD2, A_ASP_74	NH2, A_ARG_64	HH22, A_ARG_64	2.78	1.78	5.75
3IOH	OD2, A_ASP_67	NE1, A_TRP_72	HE1, A_TRP_72	2.69	1.83	24.82
3IOH	OE2, A_GLU_125	OH, A_TYR_101	HH, A_TYR_101	2.63	1.69	8.92
3IOH	OD2, A_ASP_67	NH1, A_ARG_121	HH12, A_ARG_121	2.96	2.09	25.64
3IOH	OD1, A_ASP_122	OG, A_SER_124	HG, A_SER_124	2.65	1.73	12.69

Table 10: 3IOH-specific side chain hydrogen bonding analysis with the new criteria as defined in the section of Materials and Methods. In this table, the residue naming scheme is Chain ID residue name residue number, ∠ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (∠ADH)

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
4ZGH	OE1, A_GLU_41	NE, A_ARG_44	HE, A_ARG_44	2.92	2.08	10.96
4ZGH	OE2, A_GLU_41	NH2, A_ARG_44	HH21, A_ARG_44	2.75	1.91	11.18
4ZGH	OE2, A_GLU_41	NE2, A_GLN_45	HE21, A_GLN_45	2.99	2.14	5.73
4ZGH	OD1, A_ASP_74	NE, A_ARG_64	HE, A_ARG_64	2.97	2.12	3.34
4ZGH	OD2, A_ASP_74	NH2, A_ARG_64	HH21, A_ARG_64	2.83	1.98	7.04
4ZGH	OD2, A_ASP_67	NE1, A_TRP_72	HE1, A_TRP_72	2.80	2.02	20.99
4ZGH	OE2, A_GLU_125	OH, A_TYR_101	HH, A_TYR_101	2.82	1.99	6.37
4ZGH	OD2, A_ASP_67	NH1, A_ARG_121	HH11, A_ARG_121	2.95	2.17	20.60
4ZGH	OD1, A_ASP_122	OG, A_SER_124	HG, A_SER_124	2.67	1.91	21.21
4ZGH	OE2, A_GLU_128	NZ, B_LYS_26	HZ2, B_LYS_26	2.77	2.03	27.77
4ZGH	OXT, B_GLY_37	NE1, B_TRP_31	HE1, B_TRP_31	2.92	2.09	12.63

Table 11: 4ZGH-specific side chain hydrogen bonding analysis with the new criteria as defined in the section of Materials and Methods. In this table, the residue naming scheme is Chain ID_residue name residue number, \angle ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (\angle ADH)

Here, in this article, the two experimental structures with PDB IDs: 3IOL [17,18] and 4ZGM [19,36] are both determined by X-ray crystallography, which is a non-dynamic method due to preparation of samples and crystallization. Therefore, a new criteria was also employed here for a high-throughput screening of electrostatic interactions for the two experimental structures with PDB IDs: 3IOL and 4ZGM [17-19,36].

Among the two sets of side chain hydrogen bonds at the GLP-1-GLP-1R binding interface

(Tables 12 and 13), one extraordinary interfacial side chain hydrogen bond was found to exist between the oppositely charged side chains of Ser18 of GLP-1 and Thr29 of GLP-1R for 4ZGM, as described with the details in Table 7. Of further structural and biophysical interest, three additional interfacial hydrogen bonds (according to the new criteria) are identified to be formed between B_ARG_36 of semaglutide backbone and A_GLU_68 of GLP-1R in 4ZGM, as listed in Tables 12 and 13.

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
3IOH	OG1, A_THR_29	OG, B_SER_18	HG, B_SER_18	3.26	2.42	24.61
3IOH	OE1, A_GLU_128	NZ, B_LYS_26	HZ1, B_LYS_26	3.21	2.54	41.12
3IOH	OE2, A_GLU_128	NZ, B_LYS_26	HZ1, B_LYS_26	2.92	2.28	42.31

Table 12: 3IOH-specific interfacial side chain hydrogen bonding analysis with the new criteria as defined in the section of Materials and Methods. In this table, the residue naming scheme is Chain ID_residue name residue number, \angle ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (\angle ADH).

PDB	Acceptor (A)	Donor (D)	Hydrogen (H)	D-A (Å)	H-A (Å)	∠ADH(°)
4ZGH	OE1, A_GLU_128	NZ, B_LYS_26	HZ2, B_LYS_26	3.41	2.69	31.93
4ZGH	OE2, A_GLU_128	NZ, B_LYS_26	HZ2, B_LYS_26	2.77	2.03	27.77
4ZGH	OE1, A_GLU_68	NE, B_ARG_36	HE, B_ARG_36	2.75	2.13	37.73
4ZGH	OE2, A_GLU_68	NE, B_ARG_36	HE, B_ARG_36	4.14	3.30	9.50
4ZGH	OE2, A_GLU_68	NH1, B_ARG_36	HH11, B_ARG_36	4.83	4.13	31.95

Table 13: 4ZGH-specific interfacial side chain hydrogen bonding analysis with the new criteria as defined in the section of Materials and Methods. In this table, the residue naming scheme is Chain ID_residue name residue number, \angle ADH represents the angle formed by acceptor (A), donor (D) and hydrogen (H) (\angle ADH).

5. Conclusion

With two sets of computational structural biophysical analyses of two experimental structures of the GLP-1-GLP-1R complex (PDB IDs: and 4ZGM), this article employed a high-throughput approach for the extraction of electrostatic structural features from GLP-1-GLP-1R complexes, including two sets of interfacial structure-stabilizing salt bridges and hydrogen bonds for the GLP-1-GLP-1R complex structures [17-19, 36]. These electrostatic

structural features contribute to the elucidation of the direct binding and interaction between GLP-1 and GLP-1R, providing biophysical insights into the stabilization of the GLP-1-GLP-1R complex structure and the molecular basis of GLP-1/GLP-1R signaling. Furthermore, this study underscores the significance of integrating computational approaches with experimental ligand-receptor structures to advance our understanding of protein-ligand interactions and accelerate drug discoveryefforts in the field of

diabetes and metabolic disorders [46,47].

Towards a GLP-1-GLP-1R-based mini GIBAC: A Brief Future Perspective

On August 11, 2022, the concept of a general intermolecular binding affinity calculator (GIBAC) was for the first time proposed in a preprint, which defined a collective set of the standards (as defined below) of a truly general intermolecular binding affinity calculator, i.e., a truly GIBAC:

- 1. a truly GIBAC needs to take genetic variations into account;
- 2. a truly GIBAC needs to work even without structural information;
- 3. for a truly GIBAC, a variety of factors need to be considered, such as temperature, pH [49,50], site-specific protonation states (e.g., side chain pKa of protein), post-translational modifications (PTMs, Figure 8), post-expression modifications (PEMs), buffer conditions, et cetera;
- 4. a truly GIBAC is able to be used the other way around, i.e., to be used as a search engine for therapeutic candidate(s). With such a GIBAC-based search engine, a list of therapeutic candidates can be retrieved and ranked according to drug-target Kd value(s), with input parameters including drug target(s) and a desired drug-target Kd value or a range of it [48,51,52,25,53,54,7,55,56].

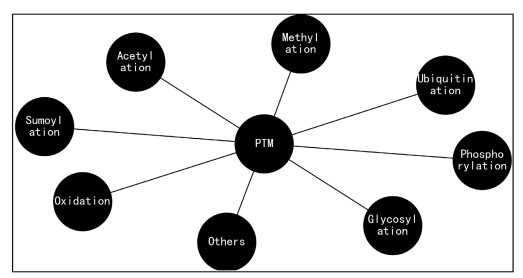


Figure 8: A range of post-translational modifications (PTMs) for biomolecules such as protein, where side chain placement and energy minimization algorithms [57] are useful to incorporate structural information of PTMs (Figure 8) and PEMs [7,55] into structural models of biomolecules such as protein, similar to the way the structure of semaglutide is modified with a C-18 fatty di-acid chain at position 26 that provides strong, specific binding to albumin [33].

As is known, the entire space of molecular types and drug modalities is vast [58], extending far beyond proteins and small molecules, which makes a comprehensive physics-based exploration practically impossible and unnecessary [59]. Nonetheless, AI algorithms rely on huge amounts of data to learn and train continuously, where its quantity and quality is inextricably linked to the performance of the model [60,61].

As charted out previously in [48], therefore, the construction of GIBAC requires two key ingredients, i.e., data and algorithm, where algorithm is like the engine of a car, and data the appropriate fuel or power source of it. Here, this article argues that in addition to data and algorithm, the construction of a real GIBAC with adequate accuracy and efficiency also requires our knowledge of biophysics underlying the structure, folding, dynamics, and the direct binding and interaction between ligand and receptor, e.g., the electrostatic structural features extracted from the two experimental structures represent the GLP-1-GLP-1R complex, with PDB IDs: 3IOL [17,18] and 4ZGM [19,36].

To sum up, the development of a GLP-1-GLP-1R-based mini GIBAC perspective represents a promising strategy for extracting

electrostatic structural features from ligand-receptor complexes and advancing drug discovery efforts targeting the GLP-1/GLP-1R axis. By combining computational modeling with experimental validation, a GLP-1-GLP-1R-based mini GIBAC offers a powerful tool for elucidating the molecular basis of ligand recognition and receptor activation, paving the way for the development of next-generation therapeutics with improved efficacy and safety for patients with diabetes and/or obesity [14–16].

Ethical Statement

No ethical approval is required.

Declaration of generative AI and AI-Assisted Technologies in The Writing Process

During the preparation of this work, the author used OpenAI's Chat GPT in order to improve the readability of the manuscript, and to make it as concise and short as possible. After using this tool, the author reviewed and edited the content as needed and takes full responsibility for the content of the publication.

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