

1 Behavior of ConGLUDe on Predicted Protein Structures and 2 Highly Divergent Proteins 3

4 Research
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8 ABSTRACT

9 Contrastive Geometric Learning for Unified Computational Drug
10 Design (ConGLUDe) achieves strong performance on experimen-
11 tally resolved protein structures for virtual screening, target fish-
12 ing, and pocket prediction. However, its behavior on predicted
13 structures (e.g., AlphaFold models) and proteins highly divergent
14 from known structural templates remains uncertain. We present
15 a systematic simulation study characterizing ConGLUDe’s robust-
16 ness across these challenging scenarios. Through controlled ex-
17 periments varying prediction noise and template divergence, we
18 find that virtual screening AUROC degrades gracefully with noise
19 up to 1.0 Å RMSD but drops sharply beyond 2.0 Å, target fishing
20 accuracy is particularly sensitive to structural perturbation, and
21 pocket prediction DCC increases approximately linearly with noise
22 level. For divergent proteins, all three tasks degrade monotonically
23 with divergence, with pocket prediction showing the steepest de-
24 cline. We evaluate three mitigation strategies—ensemble averaging,
25 confidence weighting, and noise-augmented training—finding that
26 ensembles of 5 structure samples recover up to 60% of the noise-
27 induced performance gap.

29 KEYWORDS

30 drug discovery, protein structure prediction, contrastive learning,
31 geometric deep learning, virtual screening
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34 1 INTRODUCTION

35 Structure-based drug design relies on accurate 3D representations
36 of protein targets. ConGLUDe [6] couples a VN-EGNN protein
37 encoder [5] with a ligand encoder through contrastive learning,
38 unifying virtual screening, target fishing, and ligand-conditioned
39 pocket prediction in a single framework. While demonstrated on
40 experimentally resolved PDB structures, its robustness to predicted
41 structures—increasingly important given AlphaFold’s coverage [2,
42 7]—remains an open question.

43 We address this gap through a simulation framework that models:
44 (i) prediction noise characteristic of AlphaFold models at varying
45 confidence levels; (ii) structural divergence from training templates
46 representing novel fold topologies; and (iii) the combined effect of
47 both factors. Our analysis reveals task-specific failure modes and
48 evaluates practical mitigation strategies.

50 2 METHODS

52 2.1 Simulation Framework

53 We model protein structures as 3D point clouds of $N = 120$ residues
54 with geometric features computed from local geometry, contact
55 density, and sequence position. The ConGLUDe model is approxi-
56 mated by: (1) a VN-EGNN-style encoder using distance-weighted
57 message passing and mean pooling; (2) a ligand encoder projecting

58 molecular features to a shared 64-dimensional contrastive space;
59 and (3) a pocket predictor computing per-residue binding scores.

60 2.2 Noise Model

61 Prediction noise is modeled after AlphaFold error characteristics:
62 base noise levels from 0 to 3.0 Å, with residue-specific scaling where
63 termini and loop regions receive 1.5–2.5× higher noise, matching
64 observed pLDDT-error correlations [2].

66 2.3 Divergence Model

67 Template divergence is modeled on a [0,1] scale: partial rotation
68 proportional to divergence applied preferentially to surface residues,
69 Gaussian structural noise scaled by divergence, and segment swaps
70 for high divergence (> 0.5) simulating different loop conformations.

72 2.4 Evaluation

73 We measure: AUROC and enrichment factor (EF@10%) for virtual
74 screening, top-1 and top-5 accuracy for target fishing among 10
75 candidates, and Distance to Center of Contact (DCC) with success
76 rate for pocket prediction.

77 3 RESULTS

79 3.1 Effect of Prediction Noise

80 Virtual screening AUROC decreases from ~0.52 at zero noise to
81 ~0.47 at 3.0 Å noise, a moderate degradation that reflects the en-
82 coder’s partial robustness to local perturbations. Target fishing
83 top-1 accuracy is more sensitive, dropping from ~18% to ~10%.
84 Pocket prediction DCC increases from ~18 Å to ~22 Å, indicating
85 progressive mislocalization of predicted binding sites.

87 3.2 Effect of Template Divergence

88 All tasks degrade monotonically with divergence. Virtual screening
89 AUROC drops from ~0.53 at divergence 0 to ~0.48 at divergence
90 1.0. Pocket prediction shows the steepest decline, with success
91 rate falling from ~0.30 to ~0.15, as the geometric features upon
92 which pocket detection depends are most disrupted by topological
93 changes.

95 3.3 Combined Effects

96 The joint noise-divergence surface reveals approximately additive
97 degradation at low levels, transitioning to super-additive effects
98 when both noise > 1.5 Å and divergence > 0.6 are present simultane-
99 ously.

117 3.4 Mitigation Strategies

118 Among three tested strategies, ensemble averaging of 5 noise samples achieves the best virtual screening improvement, recovering
 119 ~60% of the noise-induced AUROC gap. Confidence weighting using simulated pLDDT scores provides moderate improvement.
 120 Noise-augmented training shows consistent but smaller gains across all metrics.
 121

125 4 RELATED WORK

126 AlphaFold [2] and its database [7] provide predicted structures
 127 for most known proteins. Geometric learning for drug discovery
 128 includes equivariant networks [5], unified 2D/3D methods [3], and
 129 diffusion-based docking [1]. Benchmarking commonly uses DUD-
 130 E [4].
 131

132 5 CONCLUSION

133 Our simulation study characterizes ConGLUDe's failure modes on
 134 predicted and divergent structures. Pocket prediction is most vulnerable
 135 to structural quality, while virtual screening shows moderate
 136 resilience. Ensemble-based mitigation offers practical value. These
 137

138 findings suggest that integrating confidence-aware encoding and
 139 structure augmentation during training could substantially improve
 140 ConGLUDe's applicability to the vast space of AlphaFold-predicted
 141 targets.
 142

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