

RESEARCH ARTICLE

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Reverse Vaccinology Approach for Identification of Potential Vaccine Candidates against *Vibrio alginolyticus* using Integrated Omics Strategies

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Abstract

Vibrio alginolyticus is an opportunistic marine pathogen that causes severe vibriosis in aquatic animals and occasionally in humans, leading to substantial economic losses in aquaculture. Despite progress in antimicrobial therapy, the emergence of multidrug-resistant (MDR) strains and the lack of effective vaccines have emphasized the need for novel approaches. The objective of this study was to identify and assess vaccine candidates in *V. alginolyticus* ATCC 17749 using an integrative strategy that integrates reverse vaccinology and immune-informatics. The complete proteome of *V. alginolyticus* ATCC 17749 was retrieved from NCBI databases, and computational pipelines were used to predict subcellular localization, transmembrane topology, and antigenicity. Surface-exposed, non-allergenic and non-toxic outer membrane and secretory proteins with high antigenicity scores were selected for epitope prediction. Cytotoxic T lymphocyte (CTL), helper T lymphocyte (HTL), and B-cell epitopes were identified through NetMHCpan 4.1, IEDB, and ABCpred, respectively. Robust humoral and cellular immune responses were predicted by immune simulation. The target proteins were docked with TLR2 and TLR4 using HDock, and the simulation was performed using the iMODS server for the highest-scoring docking complex for each receptor. In silico cloning to express the target protein was performed using the SnapGene tool. This integrated computational vaccinology approach reliably identifies promising antigenic targets for *V. alginolyticus*, providing a foundation for the rational development of next-generation polyvalent vaccines against *Vibrio* infections in aquaculture and related fields.

Keywords: Reverse Vaccinology, Immune-informatics, Outer Membrane Proteins, Extracellular Proteins, Multiepitope Prediction

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INTRODUCTION

Vibrio alginolyticus, a Gram-negative, thermophilic, marine bacterium found worldwide in seawater and estuaries and is isolated frequently from shellfish and marine fish.¹ The opportunistic pathogen causes vibriosis, infecting various marine organisms, including oysters, fish, sea cucumbers, shrimp, mussels and seahorses. It can cause infection in humans, including endophthalmitis, otitis, wound infections and diarrhea. Consequently, this bacterium poses a serious zoonotic threat to both humans and animals. Several characteristics, such as biofilm development, adhesion, cytotoxicity, and the presence of a *Vibrio* pathogenicity island (VPI), are linked to *V. alginolyticus* virulence.²

The increasing emergence of lactamase, extended-spectrum β -lactamase, kanamycin, and tetracycline-resistant genes, mainly driven by horizontal gene transfer, has contributed to the rise of multidrug-resistant (MDR) strains of *V. alginolyticus*.^{3,4} As conventional antibiotics become less effective, vaccination emerges as a promising, environmentally sustainable strategy for protecting aquaculture systems against vibriosis. Although recombinant and whole-cell protein vaccines have been developed for *Vibrio* species, their efficacy is often strain-specific, short-lived, and constrained by antigenic variability.⁵ Consequently, identifying cross-protective for broad-spectrum antigens is an essential focus in vaccinology. Interpreting the mechanisms of infection is key for identifying optimal drug targets and disease control strategies, thereby preventing outbreaks in aquaculture.¹

Extensive genomic studies of *V. alginolyticus* have revealed that two circular chromosomes, totaling around 5.2-5.5 Mb, with a GC content of about 44.8%, along with several plasmid-borne virulence and resistance factors.⁶ In addition to secretion systems and stress-response components that aid host colonization, conserved virulence genes such as *toxR*, *ompU*, *vas*, *vop*, and *hlyA* have been identified by comparative genomic and pan-genomic analyses.⁷ Despite the availability of more than 70 complete and draft genomes, the immunogenic landscape of *V. alginolyticus* remains poorly characterized. The immune-protective potential of outer

membrane proteins (OMPs) like BamA, OmpK, OmpU, and LptD is conserved throughout various *Vibrio* species. It is a promising vaccine target, as highlighted by previous studies.^{8,9} Nevertheless, only a few comparable studies have focused on *V. alginolyticus*, and the essential immunodominant epitopes necessary for vaccine development remain largely unidentified.

Reverse vaccinology and immune-informatics have transformed antigen discovery by integrating genomic, proteomic, and structural data to identify potential vaccine targets and accurately predict B- and T-cell epitopes. These methods permit rapid in silico screening of entire bacterial proteomes for surface-exposed, antigenic, and non-allergenic proteins capable of eliciting protective immune responses.¹⁰ Application of these approaches to *V. alginolyticus* decreases dependence on culture-based antigen screening, addresses limitations of conventional vaccine development, and facilitates the creation of more effective and safer multi-epitope vaccines. The present study identifies and characterizes potential vaccine candidate proteins in *V. alginolyticus* by integrating reverse vaccinology and immune-informatics tools, thereby establishing a computational framework for multi-epitope vaccine design, targeting this for economic and medical significance (Figure 1).

MATERIALS AND METHODS

Proteome sequences retrieval

The nucleotide sequence (RefSeq GCF_000354175.2) of *V. alginolyticus* ATCC 17749 was retrieved from the NCBI database. This sequence was used for comprehensive computational analyses to identify potential vaccine candidate proteins and their epitopes for peptide vaccine development.

Subcellular localization

As an important step in identifying potential vaccine candidates, the subcellular localization of proteins in the genome was predicted using the independent web server PSORTb v3.0.3 (<https://www.psort.org/psortb>).¹¹ Subcellular localization is required for secreted and outer-membrane proteins, as they interact with host molecular receptors, triggering a

signal cascade that induces an adaptive immune response.¹² A total of 4,467 encoded proteins were uploaded to PSORTb server. Only proteins containing N-terminal signal peptides and secretory signals were shortlisted for DeepLocPro-1.0 (<https://services.healthtech.dtu.dk/services/DeepLocPro-1.0>) to enhance and cross-check the reliability and accuracy of predictions for proteins localized to the outer membrane or the extracellular space.^{13,14}

Signal peptide prediction

The presence of signal peptides, which facilitate translocation across membranes, was determined using SignalP 6.0 (<https://services.healthtech.dtu.dk/services/SignalP-6.0>) under the other organism and slow model, yielding long outputs.¹⁴ LipoP 1.0 (<https://services.healthtech.dtu.dk/services/LipoP-1.0>) with the extensive output format, including graphics, was used to detect lipoproteins.¹⁵

Predicting the transmembrane helices

Transmembrane topology was analyzed using DeepTMHMM (<https://dtu.biolib.com/DeepTMHMM>) validating with Phobius (<https://phobius.sbc.su.se/>) to predict α -helical transmembrane regions.^{16,17} Proteins that consisted of more than one transmembrane helix were excluded to avoid poor expression yield and folding issues in vaccine production systems.¹⁸

Antigenicity

Antigenicity evaluation was performed using Vaxijen v2.0 (<http://www.ddg-pharmfac.net/vaxijen/VaxiJen/VaxiJen.html>) with a threshold value of 0.5, optimized for bacterial models.¹⁹ It employs a partial least squares (PLS) algorithm for predicting antigens.^{19,21}

Allergenicity and toxicity analysis

To ensure safety and minimize potential hypersensitivity risks, the allergenic proteins were identified by AllerTOP v2.1 (<https://www.ddg-pharmfac.net/AllerTOP>), while toxicity was assessed using ToxinPred2 (<https://webs.iitd.edu.in/raghava/toxinpred2>) using Hybrid (RF+BLAST+MERCI) with a threshold of 0.6. Proteins classified as non-allergenic and non-toxic were shortlisted for further analysis.^{22,23}

Adhesin probability

The adhesin probability was determined using SPAAN via Vaxign, demonstrating the ability to identify adhesins at high Pad values throughout diverse bacterial species. Proteins with an adhesin probability of 0.5 or higher were classified as adhesion-related and deemed likely to interact with host cells.^{24,25}

Homology analysis

The non-homologous human host proteins of the pathogen were identified using a standalone NCBI BLASTp with Position-Specific Iterative BLAST (PSI-BLAST) against the *Homo sapiens* (taxid:9606) dataset obtained from NCBI. In the homology analysis, values $\leq 35\%$ identity and $\leq 35\%$ query coverage were used as thresholds, and the proteins to be significantly similar were not selected, while the analysis of significantly non-similar proteins was performed in the computational pipeline.^{26,27}

Physicochemical properties and conservative analysis of predicted target proteins in all *V. alginolyticus* strains

Predicted target proteins were aligned to the genomes of all known *V. alginolyticus* strains to assess protein conservation and identify broad-spectrum vaccine candidates. The tBLASTn tool (<https://blast.ncbi.nlm.nih.gov>) was employed to compare these proteins against *V. alginolyticus*, Sakazaki 1968 (taxid:663), *V. alginolyticus* 12G01 (taxid:314288), *V. alginolyticus* (chemovar. iophagus) (taxid:664), *V. alginolyticus* E0666 (taxid:1231058), *V. alginolyticus* 40B (taxid:674977), and *V. alginolyticus* NBRC 15630 (taxid:1219076).²⁸ Proteins with at least 50% pairwise identity and at least 80% query coverage were categorized as highly conserved proteins, for which ProtParam tool (<https://web.expasy.org/protparam>) depicted the stability and physiochemical properties.²⁹

Immune simulation analysis

C-immSim (<https://kraken.iac.rm.cnr.it/C-IMMSIM/index.php>) was used to evaluate the immunogenic potential of proteins. The agent-based method stimulates humoral, innate and cell-mediated immunity. In the simulation, the antigen was modified to be devoid of

lipopolysaccharide, while all other parameters were selected according to their default values. The Th-1-based immune response was indicated by the induced production of interleukins (IL-10, IL-12, and IL-2) and interferon gamma (IFN- γ), as well as by surging antibody titers of primary and secondary antibodies, such as IgM and IgG. These factors were the main components for the

assessment of protein immunogenicity. This step enables the prediction of the immunogenicity of the target proteins for multipeptide prediction.³⁰

B-cell epitopes: prediction and screening

B lymphocytes, a fundamental element of the humoral immune system, initiate and enhance the response of the adaptive immune

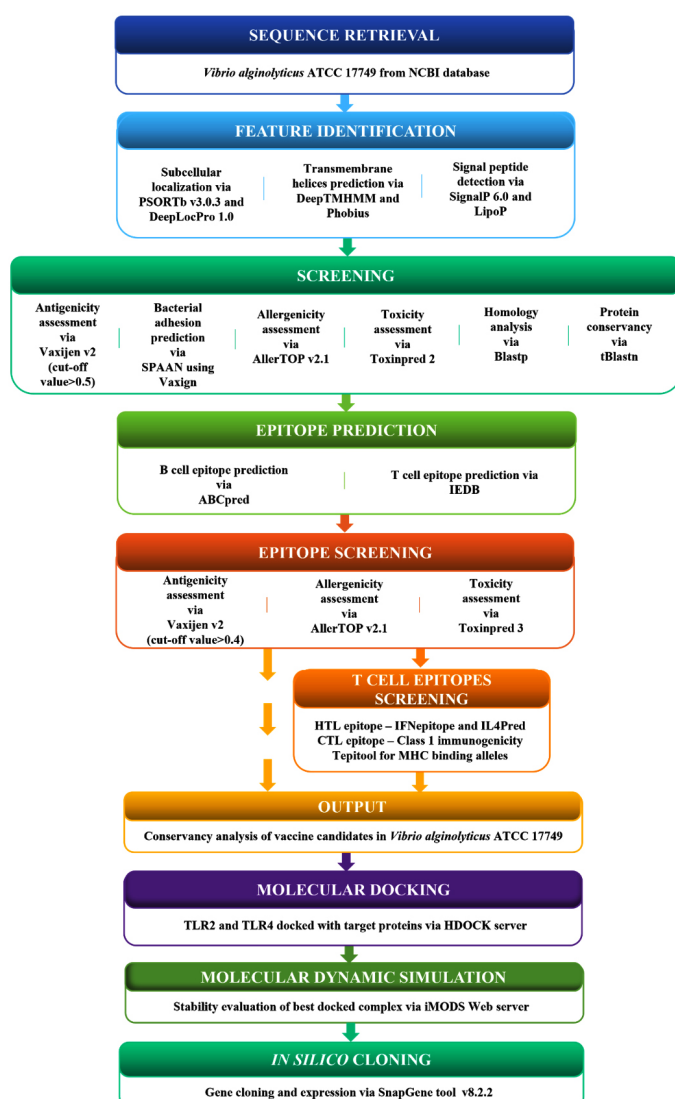


Figure 1. Workflow for computational prediction of vaccine candidate proteins and epitopes against *Vibrio alginolyticus* ATCC 17749. Protein sequences were screened for localization, transmembrane regions, signal peptides, antigenicity, adhesion, allergenicity, toxicity, homology, conservation, and physicochemical properties. B- and T-cell epitopes were predicted, followed by molecular docking with human TLR2/TLR4, simulation studies, and in silico cloning using SnapGene

system through interactions between B-cell receptors (BCRs) and B-cell receptor epitopes. ABCpred (<https://webs.iiitd.edu.in/raghava/abcpred/>) was used to predict Linear B-cell (LBL) with a 0.51 threshold and 16 amino acids as the epitope length, employing recurrent neural network algorithms. Vaxijen v2.0 servers using 0.4 as threshold was used predicted antigenicity while allergenicity and toxicity analysis with AllerTOPv2.0 (<https://www.ddg-pharmfact.net/allertop>) and ToxinPred3 (<https://webs.iiitd.edu.in/raghava/toxinpred3>), respectively, wherein non-toxic and non-allergenic epitopes were considered for conservancy analysis.

Cytotoxic T lymphocytes (CTLs) Epitopes: prediction and screening

As the antigenic peptide is presented to the T-cell receptor, the MHC molecule's interaction becomes a highly selective phase. Therefore, CTL epitopes prediction was done using the NetMHCpan 4.1 (<https://services.healthtech.dtu.dk/service.php?NetMHCpan-4.1>) server. Strong binders were identified using a threshold of 0.5%, and the epitope length was set to 9-mers. The epitopes identified as strong binders, antigenic, non-allergenic and non-toxic, similar to those assessed for B-cell epitopes, were then analyzed for immunogenicity using the Class I Immunogenicity tool in the IEDB database (<http://tools.iedb.org/>)

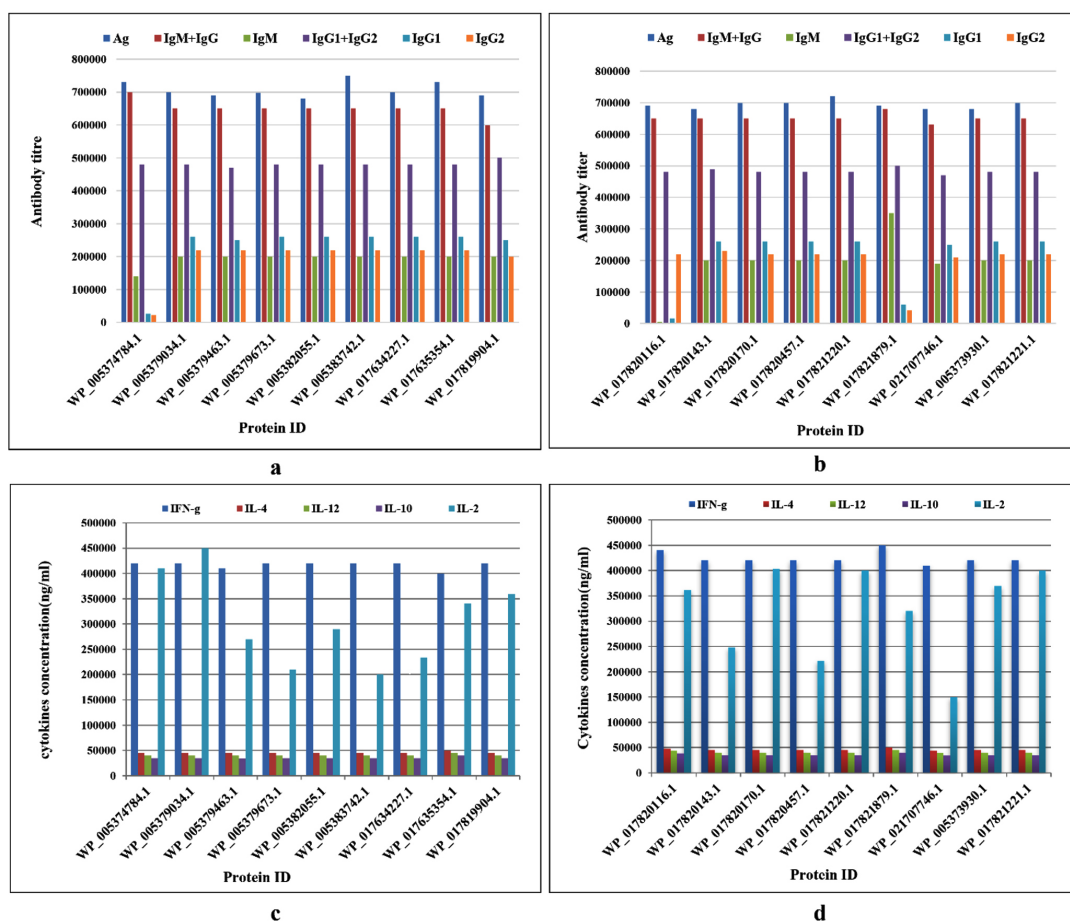


Figure 2. (a) and (b) represent antibody titers against final candidate proteins, which is measured as IgG + IgM (dark red), IgM (green), IgG1 + IgG2 (purple), IgG1 (blue), and IgG2 (orange). (c) and (d) represent induced Cytokine concentrations (ng/ml), measuring IFN- γ (dark blue), IL-4 (dark red), IL-12 (green), IL-10 (purple) and IL-2 (blue). All proteins induced strong IL-12 and IFN- γ responses, which indicates a potent Th1 (T helper) immunity

immunogenicity), retaining only scores above zero. The TepiTool feature within IEDB was also used to predict MHC class I alleles binding affinity to these epitopes. Prediction of peptides with percentile rank $\leq 1\%$ were considered as the promising candidates to develop a vaccine.

Helper T-Lymphocyte (HTL) epitopes: prediction and screening

Helper T lymphocytes (HTLs) constitute important parts of the human immune system, facilitating the activation and proliferation of cytotoxic T lymphocytes.³¹ Helper T lymphocyte (HTL) epitopes of 15-mer peptides prediction was done by IEDB MHC-II binding tool (<https://www.iedb.org>), focusing on human HLA-DR alleles with percentile ranks of $\leq 1\%$.³² The percentile rank threshold of 1% was selected in accordance with established practices to maximize the likelihood of selecting high-affinity binders with potential immunogenicity, as lower percentile ranks correlate with stronger predicted binding to MHC-II molecules. Antigenicity for predicted epitopes was set at a threshold exceeding 0.4 to maintain a stringent criterion for candidate immunogens. Epitopes identified as non-allergenic and non-toxic were further analyzed for the capacity to elicit cytokine responses. This screening was conducted using IFNepitope (<https://webs.iiitd.edu.in/raghava/ifnepitope/predict.php>) and IL-4pred (<https://webs.iiitd.edu.in/raghava/il4pred/predict.php>) for the analysis of interferon- γ and IL-4 production, respectively. TepiTool from the IEDB database was used to identify epitopes with positive scores that predict binding to MHC-II molecules, using the percentile scores below 2%.

Conservancy analysis of vaccine candidates

A conservancy analysis of the epitopes corresponding to the target proteins of *V. alginolyticus* was conducted with the IEDB Conservancy Analysis tool (<http://tools.iedb.org/conservancy>). This tool aligns predicted epitopes with homologous sequences in the target protein sourced from the NCBI database. Epitopes conserved in at least 95% of the analyzed strains were selected as the final vaccine candidates.

Molecular docking

In silico cloning ensures that a particular

host will express the vaccine protein when the gene is cloned into a suitable vector. The amino acid sequences of human TLR2 and TLR4 were retrieved from UniProt with primary accession numbers O60603 and O00206. Molecular docking was performed using the HDock server and sequences were uploaded for all the target proteins with TLR2 and TLR4, respectively.³³ The highest predicted binding free energy for both the protein-TLR2 and protein-TLR4 complexes was further used in the simulation.

Molecular dynamics simulation

Analysis of the collective motions of the complex with the highest docking score was performed using the iMODS server (<https://chaconlab.org/multiscale-simulations/imod>).^{34,35} The structural stability of the modelled complex was validated using Normal Mode Analysis (NMA). The best docking results from HDock for the TLR2 and TLR4 protein complexes, respectively, were uploaded to the iMODS server for stimulation. The iMODS analysis demonstrated that the complex is structurally sound, with limited atomic fluctuations as indicated by B-factor and deformability profiles. Flexibility was confined to non-structural loop regions, which is characteristic of stable, functionally active immune-target proteins.

In silico cloning

The SnapGene tool version 8.2.2 was used for in silico cloning. The protein-coding sequence was retrieved from the NCBI database (GeneID 75168907), and PCR was performed using primers with NcoI (5'-CCATGG-3') and BamHI (5'-GGATCC-3') restriction sites at the 5' and the 3' ends of the DNA, respectively. A restriction cloning module from SnapGene (<https://www.snapgene.com/>) was used to clone the DNA sequence located between the BamHI and NcoI restriction sites in pET-28a(+), and the recombinant plasmid was transformed into BL21(DE3).

RESULTS

Retrieval of sequence for proteome screening

The complete proteome (RefSeq GCF_000354175.2) of the *V. alginolyticus* ATCC 17749 strain comprises 4,471 annotated proteins, obtained from the National Center for

Biotechnology Information (NCBI) database in FASTA format. The complete proteins from the bacterial proteome were analyzed using various subtractive proteomics and reverse vaccinology approaches.

Predicted extracellular or outer membrane proteins

A total of 4,471 proteins in the proteome were assessed for subcellular localization by the PSORTb and DeepLocPro server. According to classifications from both servers, 172 proteins were identified as extracellular or outer membrane proteins by PSORTb, and 147 by DeepLocPro. Only proteins predicted as outer-membrane or extracellular by both servers were included in subsequent analyses.

Screening of proteins with a signal peptide

Signal peptides, which facilitate protein transport, were analyzed in 147 proteins using SignalP 6.0. Of these, 86 had general secretory pathway (Sec/spl) signal peptides, and 29 had lipoprotein (Sec/SPII) signal peptides, totalling 115 proteins with signal peptides. LipoP analysis identified WP_005375288.1 and WP_021707593.1 as cytoplasmic signal peptides. After combining results, 110 proteins with suitable signal peptides were selected.

Transmembrane alpha helix prediction

Transmembrane helicity was assessed in 110 bacterial proteins using DeepTMHMM and Phobius. Based on this analysis, 109 proteins were shortlisted for further study including 2 proteins with one and 107 proteins with no transmembrane α -helix.

Antigenicity prediction

All 109 candidate proteins were computed for antigenicity using VaxiJen v2.0, with 0.5 as the threshold. Ninety one proteins exceeded this threshold and were shortlisted for further assessment.

Allergenicity and toxicity profiling

The 91 antigenic proteins were screened for allergenicity using AllerTOP, which led to the exclusion of 14 potential allergens. Toxicity analysis

of the remaining 77 proteins with ToxinPred2 identified 17 as toxic. The final 60 non-allergenic, non-toxic proteins were considered suitable for further vaccine development.

Adhesin probability

Adhesin probabilities in SPAAN greater than 0.5 predicted by the Vaxign server, resulting in 28 shortlisted proteins for further analysis.

Homology analysis of the targeted proteins to the human proteome

All 28 adhesin proteins were examined to identify homology with the human proteome *Homo sapiens* (taxid: 9606). No significant result of similar homology with human proteins was observed, indicating a minimal risk of autoimmune responses and cross-reactivity.

Physicochemical properties of broad-spectrum proteins for a vaccine construct against all strains of *V. alginolyticus*

tBLASTn analysis of 28 proteins against all available *V. alginolyticus* strains in the NCBI database identified 19 proteins with at least 50% pairwise identity and 80% minimum query coverage. Nine non-conserved candidate proteins were excluded.

ProtParam was utilized to evaluate the physicochemical properties of the 19 proteins, yielding data on theoretical isoelectric point (pI), amino acid composition, aliphatic index, molecular weight, and GRAVY. As one of the proteins showed a positive value of GRAVY indicating hydrophobicity as was sorted out as it may complicate the production process.³⁶ The 18 selected proteins from the whole proteome as the final vaccine targets are depicted in Table 1.

Immune simulation analysis

The 18 selected proteins were evaluated for their capacity to elicit adaptive immune responses using the C-ImmSim tool. This analysis was critical to identify candidate proteins capable to induce effective humoral and cellular immunity, which includes T-cell cytokine secretion and antibody production. The results for all promising candidates are presented in Figure 2, which illustrates immunoglobulin activity

Table 1. The homology of final predicted proteins for vaccine against *V. alginolyticus* ATCC 17749

No.	Protein ID	Characteristics	No. of amino acids	Molecular weight (Da)	Theoretical PI	Aliphatic Index	GRAVY value	No. of predicted TMHs
1	WP_005373930.1	Copper resistance protein NlpE [Vibrio]	177	19226.16	3.92	75.48	-0.427	1
2	WP_005374784.1	Malto porin [Vibrio]	423	46118.68	4.52	68.97	-0.448	0
3	WP_005379034.1	DUF2860 domain-containing protein [Vibrio]	318	35721.49	4.64	78.30	-0.361	0
4	WP_005379463.1	YjbH domain-containing protein [Vibrio]	730	83247.74	4.65	64.64	-0.494	0
5	WP_005379673.1	LPS assembly protein LptD [Vibrio]	781	89277.91	4.36	74.15	-0.594	0
6	WP_005382055.1	Hypothetical protein [Vibrio]	468	51587.31	4.54	65.75	-0.432	0
7	WP_005383742.1	Porin family protein [Vibrio]	142	20304.08	3.98	74.60	-0.229	0
8	WP_017634227.1	Oligogalacturonate-specific porin KdgM family protein [Vibrio]	257	29547.77	5.65	61.05	-0.710	0
9	WP_017635354.1	AcfA family outer membrane beta-barrel protein [Vibrio]	215	23772.32	4.57	78.51	-0.256	0
10	WP_017819904.1	LruC domain-containing protein [Vibrio]	701	75540.24	4.11	71.81	-0.295	0
11	WP_017820116.1	Endonuclease [Vibrio]	538	59392.76	4.59	59.78	-0.557	0
12	WP_017820143.1	N-acetylglucosamine-binding protein GbpA [Vibrio]	487	53599.49	4.68	70.94	-0.473	0
13	WP_017820170.1	Endonuclease/exonuclease/phosphatase family protein [Vibrio alginolyticus]	287	32373.34	8.77	82.54	-0.437	0
14	WP_017820457.1	Outer membrane beta-barrel protein [Vibrio]	213	23764.44	4.41	92.91	-0.118	0
15	WP_017821220.1	Hypothetical protein [Vibrio]	581	62225.40	3.91	68.81	-0.561	0
16	WP_017821221.1	Polysaccharide lyase family 7 protein [Vibrio]	521	57628.08	4.96	64.97	-0.546	0
17	WP_017821879.1	Porin [Vibrio]	351	38697.69	4.71	77.04	-0.458	0
18	WP_021707746.1	MipA/OmpV family protein [Vibrio]	434	49532.09	4.77	74.17	-0.327	0

Table 2. List of final CTL epitopes of target proteins predicted by NetMHCpan 4.1

No.	Protein ID	CTL epitopes	MHC-I binding alleles	Antigenicity value	Allergenicity status	Toxicity status	MHC Class I immunogenicity score	Conservancy analysis ($\leq 100\%$)
1	WP 005379034.1	MLQPAFTYI	HLA-A*02:03 HLA-A*02:06 HLA-A*02:01 HLA-A*32:01 HLA-A*68:02	1.4149	-	-	0.13014	100
2	WP 005379463.1	YMPEIALGV	HLA-A*02:01 HLA-A*02:03 HLA-A*02:06 HLA-A*68:02	1.1309	-	-	0.27402	100
		KLGTDFDFTL	HLA-A*02:01 HLA-A*32:01 HLA-A*02:06	1.6785	-	-	0.30642	100
		SIPFDIMTV	HLA-A*02:06 HLA-A*68:02 HLA-A*02:03 HLA-A*02:01	1.4647	-	-	0.13556	100
3	WP 005379673.1	TLHHPRFEV	HLA-A*02:03 HLA-A*02:01 HLA-A*68:02 HLA-A*02:06	1.2583	-	-	0.23827	100
		QLDDEVSRV	HLA-A*02:01 HLA-A*02:03 HLA-A*02:06 HLA-A*01:01	0.9367	-	-	0.0565	100
4	WP 005382055.1	TLAFDFYGV	HLA-A*02:01 HLA-A*02:03 HLA-A*02:06 HLA-A*68:02	1.6693	-	-	0.27898	100
5	WP 017819904.1	IAAGIVPAV	HLA-B*51:01 HLA-A*02:06 HLA-A*68:02 HLA-A*02:01 HLA-A*02:03 HLA-B*35:01	0.4731	-	-	0.22876	100
6	WP 017820143.1	YIIESTPTT	HLA-A*02:06 HLA-A*02:01 HLA-A*02:03 HLA-A*68:02	0.4268	-	-	0.03271	100
7	WP 017820457.1	IVDENNFKI	HLA-A*02:06 HLA-A*02:01 HLA-A*01:01 HLA-A*02:06 HLA-A*02:03 HLA-A*68:02	1.1132	-	-	0.06836	100
		YVNEVGSL	HLA-A*02:01 HLA-A*26:01 HLA-B*35:01					

Table 2. Cont...

No.	Protein ID	CTL epitopes	MHC-I binding alleles	Antigenicity value	Allergenicity status	Toxicity status	MHC Class I immunogenicity score	Conservancy analysis ($\leq 100\%$)
8	WP_017821879.1	SLGNVDFSV	HLA-A*32:01	0.7135	-	-	0.06473	100
			HLA-B*08:01					
			HLA-B*07:02					
			HLA-B*53:01	0.7171	-	-	0.06771	100
			HLA-B*15:01					
			HLA-A*02:01					
			HLA-A*02:06					
			HLA-A*02:03					

and cytokine induction patterns representing immunological profiles of the final 18 target proteins, encompassing cytokine levels (IFN- γ , IL-2, IL-10, and IL-12), and antibody titers (IgG, IgM, IgG1, and IgG2).

B-Cell Epitope Prediction

Linear B-cell epitopes for 18 candidate proteins were predicted using ABCpred (threshold = 0.51), yielding 250 potential sequences. The predicted antigenic, non-allergenic, and non-toxic epitopes were analyzed for conservation across all *V. alginolyticus* strains in the NCBI database, revealing that all epitopes were $\geq 95\%$ conserved. Supplementary Table depicts the predicted LBL epitopes and their corresponding immunogenic properties.

Prediction of CTL epitopes

NetMHCpan-4.1 identified high-affinity CTL epitopes (9-mer peptides, percentile ranks $\leq 1\%$) for 18 selected proteins. T-cell epitopes that were allergenic or toxic were excluded after antigenic analysis. The remaining candidate epitopes were confirmed to be highly antigenic, non-toxic, non-allergenic, immunogenic, and conserved through comprehensive prediction and screening. 12 epitopes for 8 vaccine candidate proteins were predicted to be the final epitopes for vaccine construction. The selected CTL epitopes are shown in Table 2 with their immunogenic properties.

Prediction of HTL epitopes

HTL epitopes were predicted by the IEDB MHC-II module, yielding 48 high-confidence 15-mer peptides with percentile ranks $\leq 2\%$. The epitope analysis was the same as CTL with additional IFN-epitope and IL-4pred analyses, which further confirmed that 29 of the HTL epitopes of 12 proteins could induce IFN- γ and IL-4 cytokine responses, suggesting balanced Th1/Th2 stimulation. The final vaccine candidate epitopes are depicted in Table 3.

Conservancy analysis of the epitope

The predicted B-cell and T-cell epitopes were analyzed for their conservancy and epitopes with 100% conservancy were chosen for the final vaccine candidates.

Molecular docking

The HDock server was used to dock the protein WP_005379673 (PDB ID 4N4R) to the human immune receptors TLR2 (PDB ID 6NIG) and TLR4 (PDB ID 2Z63), and the top 10 models were selected based on docking scores. Model 1 was selected for both docking runs because it had the highest docking score. The TLR2-WP_005379673 (Figure 3a) complex and the TLR4-WP_005379673 (Figure 3b) complex had a docking score of -401.21 and -466.38, respectively. The results indicated a strong interaction between the protein and TLR2 and TLR4.

Table 3. List of selected HTL epitopes of vaccine target proteins predicted by Class I Immunogenicity tool in the IEDB database

No.	Protein ID	HTL epitopes	MHC-II binding alleles	Antigenicity value	Allergenicity status	Toxicity status	IFN- γ	IL-4	Conservancy analysis
1	WP 017820116.1	KRWHKNDPVSEKERN	HLA-DRB3*02:02	0.6209	-	-	+	Inducer	100
2	WP 017820143.1	EWTFTANHVTRDWKY	HLA-DRB1*04:05	0.8656	-	-	+	Inducer	100
			HLA-DRB3*02:02						
		FEWFTTANHVTRDWK	HLA-DRB1*04:05	1.2486	-	-	+	Inducer	100
			HLA-DRB3*02:02						
		TKPDWNPNASLSRDS	HLA-DRB3*02:02	1.0432	-	-	+	Inducer	100
			HLA-DRB5*01:01						
		WTFITANHVTRDWKY	HLA-DRB3*02:02	0.7535	-	-	+	Inducer	100
		QTFEFTTANHVTRD	HLA-DRB1*07:01	1.2071	-	-	+	Inducer	100
3	WP 017820170.1	DKPDFSITRGNSKLR	HLA-DRB5*01:01	1.2201	-	-	+	Inducer	100
4	WP 017821879.1	EGYYSQITSDKVME	HLA-DRB1*07:01						
			HLA-DRB1*09:01	0.7562	-	-	+	Inducer	100
			HLA-DRB5*01:01						
		DELEGYSQSTDKV	HLA-DRB1*15:01	0.8373	-	-	+	Inducer	100
			HLA-DRB5*01:01						
5	WP 021707746.1	YYSFKDITGERIGAG	HLA-DRB1*01:01						
			HLA-DRB1*07:01	1.2056	-	-	+	Inducer	100
			HLA-DRB1*09:01						
		YSFKDITGERIGAGL	HLA-DRB1*01:01	1.1279	-	-	+	Inducer	100
			HLA-DRB1*07:01						
6	WP 017819904.1	GKIYRVLDLSEGNAA	HLA-DRB1*04:01						
			HLA-DRB1*04:05	1.0155	-	-	+	Inducer	100
			HLA-DRB3*01:01						
			HLA-DRB3*02:02						
		DGKIYRVLDLSEGNAA	HLA-DRB1*04:01						
			HLA-DRB1*04:05						
			HLA-DRB3*01:01	1.3431	-	-	+	Inducer	100
			HLA-DRB3*02:02						
			HLA-DRB4*01:01						
		VVVKFRITETSLAGH	HLA-DRB1*04:01						
			HLA-DRB1*04:05						

Table 3. Cont...

No.	Protein ID	HTL epitopes	MHC-II binding alleles	Antigenicity value	Allergenicity status	Toxicity status	IFN- γ	IL-4	Conservancy analysis
7	WP 017634227.1	KIYRVDLSEGAANI	HLA-DRB1*07:01	0.9804	-	-	+	Inducer	100
			HLA-DRB1*08:02						
			HLA-DRB1*09:01						
			HLA-DRB3*02:02						
			HLA-DRB1*04:01						
			HLA-DRB3*01:01		-	-			
			HLA-DRB3*02:02						
8	WP 005382055.1	YAQRKISGSSKIND	HLA-DRB1*07:01	1.0828			+	Inducer	100
			HLA-DRB1*09:01						
			HLA-DRB1*13:02						
			HLA-DRB1*15:01		-	-			
			HLA-DRB3*02:02						
			HLA-DRB3*01:01						
			HLA-DRB3*02:02						
9	WP 005379673.1	DYESEYAVNATVSYS	HLA-DRB1*15:01	0.7133			+	Inducer	100
			HLA-DRB3*02:02						
			HLA-DRB1*15:01						
			HLA-DRB5*01:01						
			HLA-DRB3*02:02						
			HLA-DRB1*04:01						
			HLA-DRB1*04:05						
10	WP 005379463.1	ASGQYYDLNENADL	HLA-DRB3*01:01	0.5506	-	-	+	Inducer	100
			HLA-DRB3*02:02						
			HLA-DRB1*04:01						
			HLA-DRB1*04:05						
			HLA-DRB3*01:01						
			HLA-DRB3*02:02						
			HLA-DRB1*04:01						
11	WP 005379034.1	GQYYDLNENADLEW	HLA-DRB1*04:05	0.5471	-	-	+	Inducer	100
			HLA-DRB3*01:01						
			HLA-DRB3*02:02						
			HLA-DRB4*01:01						
			HLA-DRB1*03:01						
			HLA-DRB3*01:01						
			HLA-DRB4*01:01						
11	WP 005379034.1	EAPSLQPSQSDFGGV	HLA-DRB1*03:01	0.7498	-	-	+	Inducer	100
			HLA-DRB1*03:01						
			HLA-DRB3*01:01						
			HLA-DRB4*01:01						
			HLA-DRB1*03:01						
			HLA-DRB3*01:01						
			HLA-DRB4*01:01						
11	WP 005379034.1	TDRSLARDANLYYVK	HLA-DRB3*01:01	0.5045	-	-	+	Inducer	100
			HLA-DRB3*01:01						
			HLA-DRB3*01:01						
			HLA-DRB3*01:01						
			HLA-DRB3*01:01						
			HLA-DRB3*01:01						
			HLA-DRB3*01:01						

Table 3. Cont...

No.	Protein ID	HTL epitopes	MHC-II binding	Antigenicity value	Allergenicity status	Toxicity status	IFN- γ	IL-4	Conservancy analysis
12	WP 005374784.1	PEIRVASYLTADKE RPEIRVASYLTADK	HLA-DRB1*15:01	0.6689	-	-	+	Inducer	100
			HLA-DRB1*13:02	0.6811	-	-	+	Inducer	100
			HLA-DRB1*15:01						
		FWARPEIRVASYLT ISDFYWNISGAGAG	HLA-DRB1*15:01	0.5231	-	-	+	Inducer	100
			HLA-DRB3*02:02	1.5003	-	-	+	Inducer	100

Molecular dynamics simulation

We employed molecular dynamics simulations using the iMODS web server to investigate the stability and dynamics of the top-docked protein-TLR2 and protein-TLR4 complexes. Figures 4a and 5a present the motion trajectories of the TLR2-WP005379673 and TLR4-WP005379673 complexes. The deformability plots (Figures 4b and 5b) indicate minimal structural distortion in both complexes. B-factor plots (Figures 4c and 5c) demonstrate the relationship between the mobility of the docked complex, as assessed by normal mode analysis (NMA), and the PDB score, which corresponds to the mean root mean square deviation (RMSD). The eigenvalue quantifies the energy required to deform the structure; lower eigenvalues indicate that carbon alpha atoms are more readily deformed. The eigenvalues for the TLR2-WP005379673 and TLR4-WP005379673 complexes are 1.034783e-05 (Figure 4d) and 1.947305e-05 (Figure 5d), respectively, suggesting high structural stability. Each normal mode is associated with variance plots that display both individual (purple) and cumulative (green) variance (Figures 4e and 5e). Covariance plots characterize the movement of correlated (red), non-correlated (white), and anti-correlated (blue) atoms within the dynamic regions of the complexes (Figures 4f and 5f). We further assessed the stiffness of the TLR2-WP005379673 and TLR4-WP005379673 complexes using the elastic network model. In Figures 4g and 5g, darker grey regions denote higher stiffness, while lighter areas indicate greater flexibility.

In silico cloning

The protein-coding gene (GeneID = 75168907) on chromosome 2 of *Vibrio alginolyticus* ATCC 17749 was amplified by PCR using forward and reverse primers with NcoI and BamHI restriction sites, respectively (Figure 6a). The PCR product was cloned to pet-28a (+) and transformed to BL21(DE3) (Figure 6b). The translations and ORF2 indicated a translated protein of 89.3 kDa and 781 amino acids, similar to WP_005379673, confirming expression.

DISCUSSION

Vibriosis caused by *V. alginolyticus* is increasing in prevalence, and drug-resistant strains are spreading rapidly, posing significant risks to aquaculture and public health. Although antibiotics and some experimental vaccines are available, the lack of broadly protective, durable immunization strategies remains a major challenge.³⁷ In this context, reverse vaccinology integrated with immune-informatics offers a robust alternative to conventional antigen discovery by enabling systematic, genome-wide identification of conserved and immunogenic vaccine targets.³⁸ This study represents a comprehensive computational effort to elucidate the immunogenic landscape of *V. alginolyticus* ATCC 17749 and to prioritize protective epitopes suitable for next-generation vaccine development. A key strength of this work is the initial genome-wide screening of the complete *V. alginolyticus* proteome, which facilitated unbiased identification of candidate antigens. By combining multiple subcellular localization tools, the analysis focused on plasma membrane and secretory proteins, which are readily accessible to host immune surveillance and are frequently immunodominant antigens. The convergence of PSORTb and DeepLocPro predictions substantially increased confidence in protein localization and reduced false positives, an approach recognized as best practice in computational vaccinology.

The selected proteins were surface-exposed, possessed signal peptides and contained fewer than two transmembrane helices, supporting their suitability for experimental validation.

Subsequently, filtering proteins based on antigenicity, allergenicity, toxicity, adhesion probability, host homology and protein conservancy dramatically reduced the candidate pool, highlighting the efficiency of integrative reverse vaccinology pipelines. As an outcome, Figure 7 depicts 18 non-host homologous, non-toxic, non-allergenic, antigenic, and adhesion-associated outer membrane and extracellular proteins retained from an initial proteome of 4,471 proteins.

Notably, a significant number of the shortlisted proteins, such as Maltoporin, DUF2860 domain-containing protein, MULTISPECIES: YjbH domain-containing protein, LPS assembly protein LptD, porin family protein, oligogalacturonate-specific porin KdgM family protein, AcfA family outer membrane beta-barrel protein, porin and MipA/OmpV family protein, are biologically essential β -barrel proteins that play important roles in membrane assembly, nutrient uptake, secretion systems, and virulence control. These proteins are highly conserved among *Vibrio alginolyticus* strains, as shown by conservation analysis, and thus are less likely to undergo antigenic drift. These findings are consistent with previous studies on *V. parahaemolyticus* and other

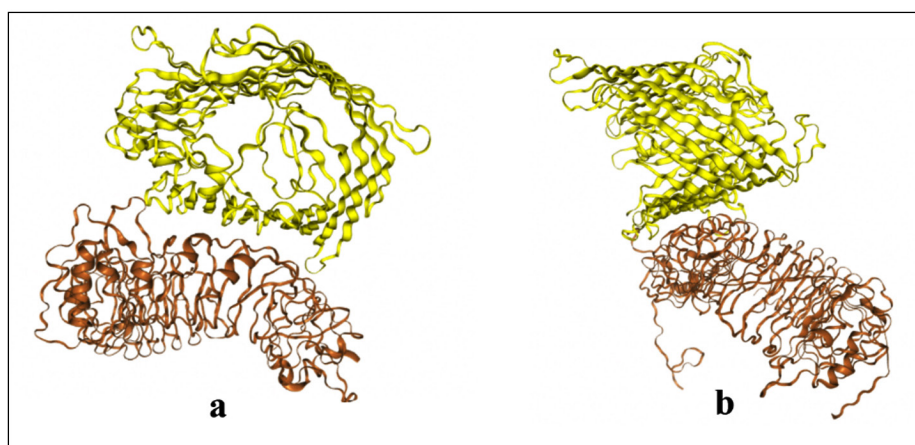


Figure 3. Molecular docking between the target protein and TLR2 (a) and TLR4 (b). The binding mode between the WP_005379673 and TLR2 and TLR4. Brown indicates TLR2 (a) or TLR4 (b), while yellow indicates the target protein

Gram-negative bacteria, in which conserved outer membrane proteins are used as effective vaccine targets.⁸ The identification of such proteins in *V. alginolyticus* provides further impetus for the development of cross-protective vaccines that can target multiple strains or species.

Immune simulation studies further confirmed the immunogenicity of the shortlisted

candidates. Several proteins were found to trigger a robust humoral and cellular immune response, characterized by higher titers of IgM and IgG, increased memory B-cell counts, and persistent cytokine secretion (Figure 2). Notably, all selected target proteins triggered robust IFN- γ and IL-12 secretion, a hallmark of the Th1-type immune response. This type of immune response

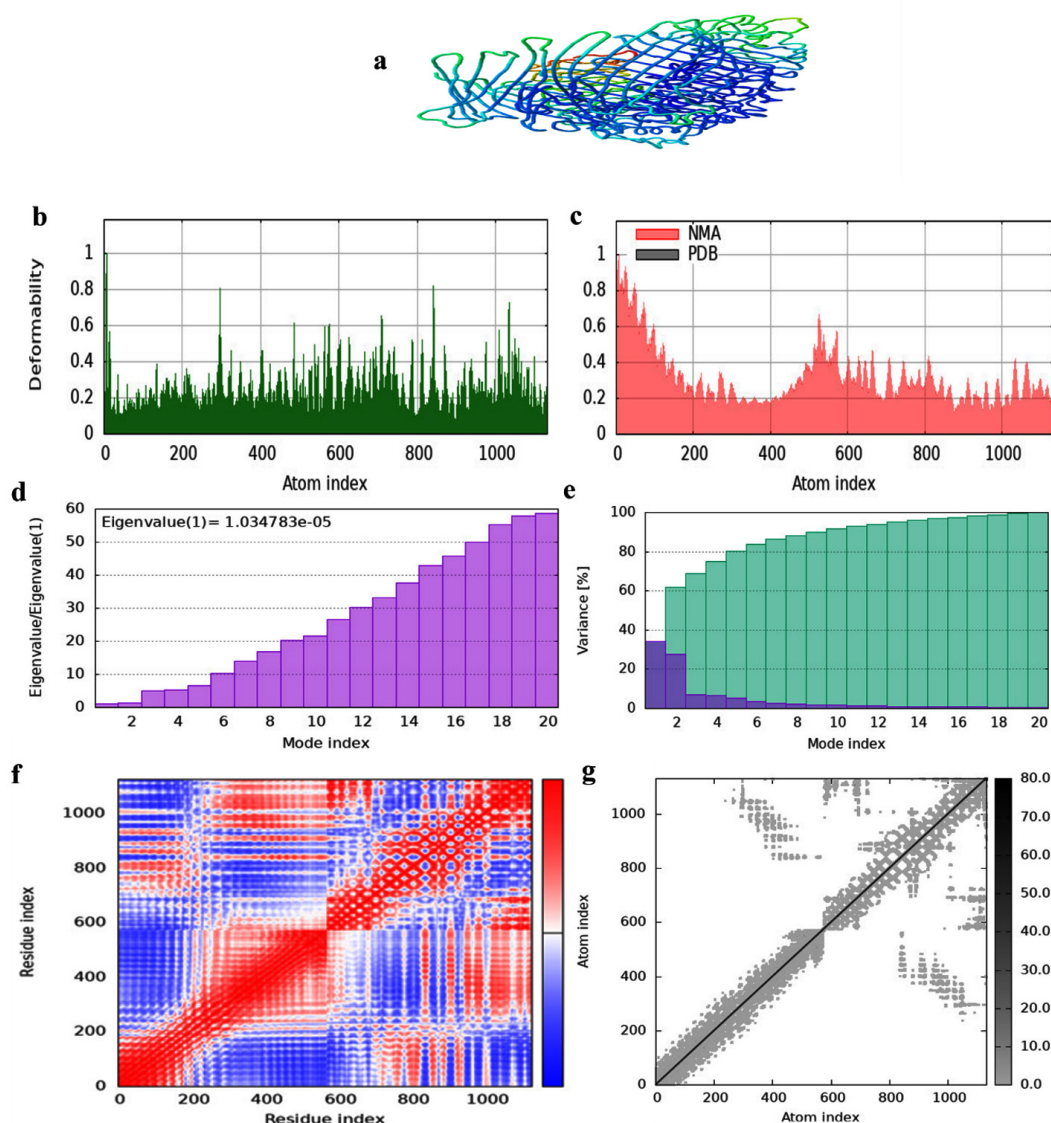


Figure 4. Molecular dynamics simulation of target protein-TLR2 complex.

(a) Mobility; (b) Deformability plot; (c) B-factor plot; (d) Eigenvalues plot; (e) Variance plot. Purple represents individual variance, while green represents cumulative variance; (f) Covariance plot. Red represents correlated motion, white represents non-correlated motion, and blue represents anti-correlated motion; (g) Elastic network. Darker grey represents stiffer regions

is especially desirable for combating extracellular bacterial infections, where it can trigger efficient macrophage activation and establishment of immune memory.

The epitope-mapping strategy used in this study further underscores the utility of immune-informatics tools in vaccine design. B-cell and T-cell epitope predictions identified several

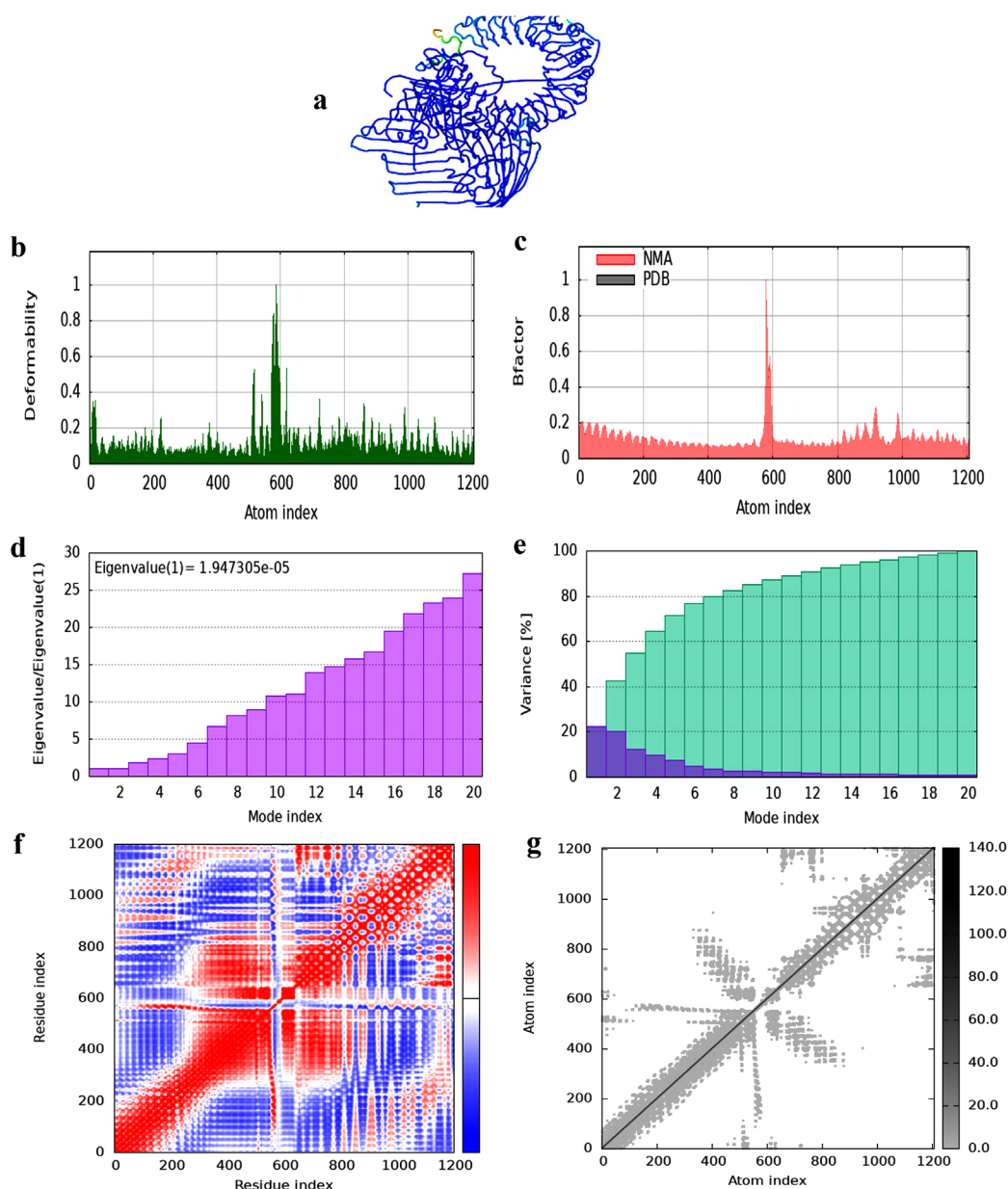


Figure 5. Molecular dynamics simulation of target protein-TLR4 complex

(a) Mobility; (b) Deformability plot; (c) B-factor plot; (d) Eigenvalues plot; (e) Variance plot. Purple represents individual variance, while green represents cumulative variance; (f) Covariance plot. Red represents correlated motion, white represents non-correlated motion, and blue represents anti-correlated motion; (g) Elastic network. Darker grey represents stiffer regions



Figure 7. Protein sorting results for the selection of the vaccine target proteins. The proteome consisted of 4,467 proteins, which underwent a computational pipeline resulting in 18 selected proteins for vaccine targets

broader, more long-lasting protection than current whole-cell or single-antigen vaccines.³⁹

This study is a strong example of how multiple fields, such as genomics, proteomics, and immunology, can be computationally integrated to address biological questions. Reverse vaccinology frameworks have been successfully applied to other pathogens such as *Pseudomonas aeruginosa*, *Klebsiella pneumoniae*, *Salmonella Typhimurium* and *Campylobacter jejuni* where large proteomes are efficiently reduced to a small number of high-confidence vaccine candidates. This work aims to apply these ideas to *V. alginolyticus*, a relatively uncharacterized but increasingly important aquaculture pathogen.⁴⁰⁻⁴³ The gradual improvement demonstrated how quickly the computational methods, which would otherwise necessitate thorough laboratory screening, could prioritize desired proteins.

Toll-like receptors (TLRs) are essential to the innate immune response, as they recognize conserved pathogen-associated molecular patterns (PAMPs) from various microorganisms.⁴⁴ To analyze their interactions, proteins were docked to TLR2 and TLR4. The highest-scoring complex, involving the protein WP_005379673, was further assessed using molecular dynamics (MD) simulations to

evaluate its stability. MD simulations utilizing normal modes from the iMODS server examined the essential dynamics and structural stability of the protein complex. Results showed no significant atomic distortion, indicating a low likelihood of deformability and a suitable level of structural rigidity. Although immune simulation and epitope prediction are useful tools for gaining insights into immunogenic potential, validation is always necessary, including recombinant expression of selected antigens, in vivo immunization trials, and evaluation of protective efficacy in appropriate animal or aquaculture models. The use of appropriate adjuvants and delivery systems is crucial to translate these predictions into effective vaccines.⁴⁵

The subunit and whole-cell inactivated vaccine against *Vibrio* infections is traditionally used, providing serotype-specific and short-term protection. Also, a vaccine can achieve higher potency if the predicted epitopes are conjugated to the cholera toxin subunit B as an adjuvant.⁴⁶ Thus, a peptide vaccine comprising immunogenic epitopes rather than the whole pathogenic 3D structure is more efficient, safer, and more convenient than the current vaccines.⁴¹

CONCLUSION

This research provides in silico predictions and assessment of potential vaccine candidate proteins and their epitopes in *Vibrio alginolyticus* ATCC 17749 using reverse vaccinology and an immune-informatics approach. The approach was successful in evaluating the proteome of the bacteria and all the strains to identify conserved, antigenic, and non-allergenic extracellular and outer membrane proteins, along with their corresponding B- and T-cell epitopes, suitable for the design of multi-epitope-based vaccines. The results clearly foreground the significant role of computational vaccinology in developing next-generation vaccines for *Vibrio* infections, especially in the wake of antibiotic resistance and climate change-driven disease emergence in aquaculture, aligning well with the systems-level and integrative focus of OMICS-driven research.

SUPPLEMENTARY INFORMATION

Supplementary information accompanies this article at <https://doi.org/10.22207/IPAM.20.3.10>

Additional file: Table.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

AUTHORS' CONTRIBUTION

AP conceptualized the study. YK performed the investigation. YK, AS and VR conducted formal analysis. YK wrote the manuscript. AP, YK, AS and VR reviewed and revised the manuscript. All authors read and approved the final manuscript for publication.

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None.

DATA AVAILABILITY

The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.

ETHICS STATEMENT

Not applicable.

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