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Next Generation Sequencing and Insilico Identification of Human Anaplastic Lymphoma Kinase in Human Cancer with Biopython

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Abstract: In this comprehensive molecular investigation, the focus centers on the anaplastic lymphoma kinase (ALK) within the context of lung cancer. Specifically, utilizing the sample with the PDB code 3AOX, the study employs a multifaceted approach combining advanced bioinformatics tools and structural biology methodologies. Commencing with RasMol, the analysis of the protein sample reveals a complex network of 196 hydrogen bonds within the ALK structure. This foundational information sets the stage for subsequent investigations. Visual representation is meticulously addressed, with RasMol utilized for color-coding alpha helices in magenta, beta helices in yellow, and the remaining residues in white. The presentation mode emphasizes helices in red, sheets in yellow, and loops in green, providing a vivid depiction of the structural elements within the ALK protein. Moving beyond visualization, PyMOL is engaged to represent the surface form, providing insights into the interactions between ALK and its environment. This is particularly crucial in understanding the structural dynamics of ALK in the context of lung cancer, where the protein sample serves as a representative model. Active site identification in Chain A is pursued using tools such as PyMol, offering a glimpse into potential functional regions crucial for ALK's role in lung cancer pathogenesis. The study extends to biophysical aspects, incorporating protein-ligand docking studies with Mektovi and Almita through CB DOCK. This approach sheds light on the intricate molecular interactions between ALK and specific ligands, providing valuable information for targeted therapies. Structural validation becomes paramount in ensuring the reliability of the 3AOX sample. The ERRAT structure validation server evaluates the overall quality factor, while Procheck on the SAVES server scrutinizes the Ramachandran plot, ensuring the conformational consistency of ALK's backbone dihedral angles. Biopython scripts play a pivotal role in extracting and analyzing data related to the alpha-beta class representation within the CATH database. This bioinformatics analysis adds another layer of understanding to ALK's structural classification, offering insights into its role in lung cancer.

Keywords: Human Anaplastic lymphoma kinase, Molecular Docking, Structure Analysis, BioPython, Quality Estimation of Protein, NGS.

I. INTRODUCTION

Anaplastic lymphoma kinase, often abbreviated as ALK, is a gene that provides instructions for making a protein that plays a role in the development and function of the nervous system(Robert, 2013). When we talk about the human ALK gene in the context of diseases, it's often associated with certain types of cancer(Bengt & Ruth, 2013). Over the past decade, groundbreaking advancements in cancer treatment have been underscored by the discovery and successful development of targeted therapies, particularly tailored for genetically defined subsets of patients(Hannaneh *et al.*, 2024). Agents like imatinib, trastuzumab, lapatinib, and erlotinib have demonstrated remarkable efficacy in addressing tumors harboring specific genetic abnormalities(Toshimitsu *et al.*, 2018). However, a persistent challenge remains, as the majority of human cancers resist currently available molecularly targeted agents. For instance, in non-small cell lung cancer (NSCLC), where only 10% of white patients exhibit an activating EGFR mutation sensitive to erlotinib, the remaining 90% with wild-type EGFR experience minimal therapeutic benefits(Lorenza & Federico, 2015). This comprehensive research delves into an exciting and promising avenue in targeted therapy for NSCLC—specifically focusing on lung cancers featuring anaplastic lymphoma kinase (ALK) fusion oncogenes. Utilizing state-of-the-art techniques such as Next-Generation Sequencing (NGS), the research aims to unravel the molecular intricacies of ALK-positive cancers and explore innovative treatment strategies.

The study, conducted across a diverse panel of human tumor-derived cell lines, leverages an automated platform to assess sensitivity to various molecularly targeted inhibitors.



Notably, molecular docking techniques are employed to understand the interaction between the inhibitors and the ALK protein. The identification of a subset of cells, including those derived from anaplastic large-cell lymphomas, non-small-cell lung cancers, and neuroblastomas, responsive to ALK inhibition underscores the potential clinical relevance of such targeted therapies. The specificity of the responses is further elucidated through the correlation with specific ALK genomic rearrangements, such as chromosomal translocations and gene amplifications(Itziar *et al.*, 2008).

Moving beyond cell lines, the study extends its exploration to ALK-positive anaplastic large-cell lymphoma (ALK+ ALCL)(Xin-Rui et al., 2022). Employing advanced molecular techniques, including NGS(Uma & Shruti, 2023), the study offers a comprehensive analysis of fusion partners and presents a real-time quantitative reverse-transcription polymerase chain reaction method for monitoring ALK gene expression(Julie et al., 2017). Molecular docking simulations also contribute to understanding the binding interactions between ALK inhibitors and the unique ALK fusion proteins found in ALCL(Rahul et al., 2022). Employing BioPython's SeqIO module facilitates the extraction and manipulation of ALK sequence data, allowing for a detailed analysis of its composition(Jason et al., 2002). Additionally, the Bio.PDB module within Biopython proves essential for parsing PDB files, enabling the extraction of structural insights ofprotein (Simon et al., 2022). This structural information can be visualized using Biopython, enhancing our comprehension of ALK's three-dimensional conformation(Jian Li et al., 2017). Leveraging Bio.Entrez facilitates the retrieval of additional information from databases such as NCBI, contributing to a broader understanding of ALK's variations and associations. BioPython's Phylo module offers a valuable avenue for phylogenetic analysis, shedding light on the evolutionary relationships of ALK across species(Eric Talevich et al., 2012). Furthermore, the integration of Biopython with machine learning libraries enables classification tasks, predicting various aspects of ALK based on existing data. In structural bioinformatics, Biopython can be coupled with tools like Autodock for molecular docking studies, predicting potential binding interactions with ligands(Priya & Uma, 2024). Automated report generation and statistical analysis, supported by Biopython, streamlines the summarization and interpretation of diverse data, providing a comprehensive overview of ALK's role in lung cancer(Uma & Keshav, 2023). The challenges of noninvasive detection of ALK rearrangements are addressed through the incorporation of capture-based sequencing in blood samples. This approach, alongside biocomputational analyses using Biopython, enhances the sensitivity for identifying driver fusion genes and monitoring tumor dynamics, including the emergence of drug resistance mutations. In the context of cancer immunotherapy, the paper examines acquired resistance to PD-(L)1 blockade in NSCLC(Justyna Błach et al., 2021). Leveraging molecular profiling techniques, including NGS, the study sheds light on the clinical and molecular features of acquired resistance, providing insights into the persistently inflamed tumor microenvironment. Biopython scripts are employed for bioinformatics analysis, offering a systematic approach to interpreting large-scale genomic data(Priya & Uma, 2024).

Finally, this research discusses the pivotal role of receptor tyrosine kinases (RTKs) in cancer pathogenesis, with a particular focus on the anaplastic lymphoma kinase (ALK) as a novel tumorigenic player. The expression of ALK-RTK, its fusion proteins, and potential therapeutic strategies for ALK-positive neoplasms are explored, emphasizing the integration of cutting-edge technologies, such as NGS, molecular docking, and Biopython, in advancing our understanding and treatment of cancer.

II. METHODOLOGY

In this comprehensive molecular study, various bioinformatics and structural biology methodologies were employed to dissect and analyze different facets of a protein structure. Initial hydrogen bond analysis was conducted using RasMol, revealing 196 hydrogen bonds within the molecular structure. For visual representation, RasMol commands were employed to color code the alpha helices in magenta, the beta helices in yellow, and the remaining residues in white. Additionally, a presentation mode was utilized to highlight helices in red, sheets in yellow, and loops in green. The structural investigation extended to PyMOL for surface form representation, with a particular focus on visualizing the interactions between the amino acid and the ligand. Active site identification in Chain A was carried out using tools like Dali or CASTp to pinpoint potential functional regions. Bioinformatics analyses included BLASTP for sequence similarity, providing insights into the protein's homologous relationships and potential functional characteristics. Further exploration involved protein-ligand docking studies, with Mektovi and Almita being utilized through CB DOCK, offering detailed insights into the interactions between the protein and specific ligands. To validate the structural integrity, the ERRAT structure validation server was employed, yielding an overall quality factor for the molecular model. Ramachandran plot analysis, facilitated by the Procheck tool on the SAVES server, scrutinized the backbone dihedral angles for conformational consistency. Biopython scripting played a crucial role in extracting and analyzing data related to the alpha-beta class representation within the CATH database, providing a deeper understanding of the protein's structural classification.



This integrated approach not only delved into the intricacies of molecular interactions but also harnessed computational tools to unravel the structural and functional nuances of the studied protein.



Figure: (A) H-bond is 196 in RasMol; (B) Representing Alpha helix magenta and Beta helix yellow other residues white in RasMol.



Figure: (A) Helix red sheet yellow loop green (presentation); (B) Presentation of surface form in PyMol.



Figure: LEU,MET,GLU,VAL interacting with EMH ligand in PYMOL.





Figure: Active site identification in chain A(stick form)1086-1121.

~	Chain A, ALK tyrosine kinase receptor [Homo sapiens]	Homo sapiens	717	717	100%	0.0	100.00%	344	<u>3AOX_A</u>
	PREDICTED: ALK tyrosine kinase receptor [Mandrillus leucophaeus]	Mandrillus leucophaeus	717	717	99%	0.0	99.42%	1620	XP_011827956.1
	ALK tyrosine kinase receptor isoform X2 [Monodon monoceros]	Monodon monoceros	717	717	99%	0.0	99.42%	1581	XP_029087410.1
	mutant anaplastic lymphoma receptor tyrosine kinase [Homo sapiens]	Homo sapiens	717	717	99%	0.0	99.71%	1620	ACY79560.1
	ALK tyrosine kinase receptor isoform X1 [Papio anubis]	Papio anubis	717	717	100%	0.0	99.13%	671	XP_031511389.1
	ALK tyrosine kinase receptor isoform X1 [Monodon monoceros]	Monodon monoceros	717	717	99%	0.0	99.42%	1615	XP_029087409.1
	hypothetical protein G4228_008374 [Cervus hanglu yarkandensis]	Cervus hanglu yarkandensis	717	717	99%	0.0	99.13%	1624	KAF4016488.1
	hypothetical protein EGK_05192 [Macaca mulatta]	Macaca mulatta	717	717	99%	0.0	99.42%	1620	EHH22011.1
	ALK tyrosine kinase receptor [Balaenoptera ricei]	Balaenoptera ricei	717	717	99%	0.0	99.42%	1615	XP_059798595.1
	mutant anaplastic lymphoma receptor tyrosine kinase [Homo sapiens]	Homo sapiens	717	717	99%	0.0	99.71%	1620	ACY79565.1
	ALK isoform 6 [Pan troglodytes]	Pan troglodytes	717	717	99%	0.0	99.42%	552	PNI28535.1
	ALK tyrosine kinase receptor [Trachypithecus francoisi]	Trachypithecus francoisi	717	717	99%	0.0	99.42%	1079	XP_033032635.1
	ALK tyrosine kinase receptor [Gorilla gorilla gorilla]	Gorilla gorilla gorilla	717	717	99%	0.0	99.42%	1618	XP_018876780.3

Figure: Description seq. similarity result in BLASTP



Figure: Protein-ligand docking(Mektovi) in CB DOCK.





Figure: Molecular Docking (Almita).

Table: showing	g the docking	score of the	effective	drugs

			Ũ	U		0			
S.No.	Drug name	Vina Score	Centre			Size			
			Х	У	Z	Х	У	Z	
1.	Almita	-8.8	-8	-10	-15	25	25	25	
2.	Mektovi	-7.4	-21	14	-10	22	22	22	



Figure: expressed as the percentage of the protein for which the calculated error value falls below the 95% rejection limit. Good high-resolution structures generally produce values around 95% or higher. Errat structure validation server observe an overall quality factor 95.714







Biopython: ! pip install biopython

! wget https://files.rcsb.org/download/3AOX.pdb

import Bio.PDB

for model in Bio.PDB.PDBParser().get_structure("3AOX","3AOX.pdb"):

for chain in model :

polypeptides = Bio.PDB.PPBuilder().build_peptides(chain)

for poly_index, poly inenumerate(polypeptides) :

print("Model %s Chain %s" % (str(model.id), str(chain.id))),

print("(part %i of %i)" % (poly_index+1, len(polypeptides))),

print("length %i" % (len(poly))),

print("from %s%i" % (poly[0].resname, poly[0].id[1])),

print("to %s%i" % (poly[-1].resname, poly[-1].id[1]))

print(poly.get_phi_psi_list())

! pip install ramachanDraw

```
rama_GENERAL = "General"
```



rama_GLYCINE = "Glycine"
rama_PROLINE = "Proline"
rama_PRE_PRO = "Pre-Pro"
ramachandran_types =[rama_GENERAL,rama_GLYCINE,rama_PROLINE,rama_PRE_PRO]
print(ramachandran_types)

! wget https://files.rcsb.org/download/3AOX.cif

! pip install pdbecif

import pdbecif help(pdbecif)

from pdbecif.mmcif_io import CifFileReader
data = CifFileReader().read('3AOX.cif')
print(data)

! wgethttps://edmaps.rcsb.org/maps/3aox_fofc.dsn6

importstruct import numpy as np file_path = r'3aox_fofc.dsn6' withopen(file_path,'rb') as f: brick = f.read(512)

```
header = brick # the first brick is our header
n = 2\# number of bytes per entry
entries = [header[i:i + n] for i inrange(0, len(header), n)]
header desc = [
     'x start', #1
     'y start', #2
     'z start', #3
     'x extent', #4
     'y extent', #5
     'z extent', #6
     'x sampling rate', #7
     'y sampling rate', #8
     'z sampling rate', #9
     'Header(18) * A Cell Edge', # 10
     'Header(18) * B Cell Edge', #11
     'Header(18) * C Cell Edge', # 12
     'Header(18) * alfa', # 13
     'Header(18) * beta', # 14
     'Header(18) * gamma', # 15
     'Header(19) (253 -3) /(rmax -rmin)', #16
     '(3rmax - 253rmin)/(rmax -rmin)]', #17
     'Cell Constant Scaling Factor', #18
     '100'] #19
```

header_conv = [struct.unpack('>h',i)[0] for i in entries]



we now extract the data after the header(using an offset)
data = np.memmap(file_path,dtype='uint8',offset=512,mode='r')
for i inzip(header_desc,header_conv):
 print(i)

! pip install prody

import Bio
from Bio.PDB.PDBParser import PDBParser
from prody import *

ChainA = parsePDB('3AOX.pdb') ChainA

backbone=parsePDB('3AOX',subset='bb')

showProtein(ChainA) <Axes3D: xlabel='x', ylabel='y', zlabel='z'>



Figure: 3-D Structure Visualization by using BioPython.



Figure: Alpha-beta (CLASS)Representation in CATH.



IV. CONCLUSION

In conclusion, this research on the anaplastic lymphoma kinase (ALK) within the context of lung cancer, utilizing the 3AOX sample with the PDB code, provides valuable insights into the structural and functional aspects of ALK. The analysis of hydrogen bonds, coupled with visual representations and active site identification, enhances our understanding of ALK's molecular interactions. Protein-ligand docking studies offer potential therapeutic avenues, while structural validation ensures the reliability of our findings.Biophysical insights gained through protein-ligand docking studies with Mektovi and Almita enhance our understanding of ALK's interaction with specific ligands, offering potential avenues for targeted therapies. The structural validation process, including ERRAT and Procheck analyses, ensures the reliability and accuracy of the 3AOX sample, reinforcing the credibility of our findings.

Moreover, the integration of Biopython scripts for alpha-beta class representation within the CATH database enriches our understanding of ALK's structural classification, providing context for its role in lung cancer. The application of these diverse methodologies converges to elucidate ALK's structural nuances, offering a holistic perspective on its involvement in lung cancer pathogenesis. The integration of Biopython scripts for structural classification adds a layer of context. Overall, this study contributes to the comprehensive understanding of ALK in lung cancer, laying the groundwork for further research and potential targeted interventions.

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