BIOMEDICAL SCIENCES

Concatenation of molecular docking and dynamics simulation of human papillomavirus type 16 E7 oncoprotein targeted ligands: In quest of cervical cancer’s treatment

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Abstract: The Human papillomaviruses type 16 E7 oncoprotein is a 98-amino-acid, 11-kilodalton acidic oncoprotein with three conserved portions. Due to its interaction with the pRb-E2F complex, CKII, CKI (mostly p21), and even HDAC1, it possesses strong transformative and carcinogenic qualities that inhibit normal differentiation and cell cycle regulation. Here, we target the E7 oncoprotein using two prior research active compounds: asarinin and thiazolo[3,2-a]benzimidazole-3(2H)-one,2-(2-fluorobenzylideno)-7,8-dimethyl (thiazolo), and valproic acid as a control. We are performing molecular docking followed by molecular dynamic analysis. By acting as competitive inhibitors in the binding site, it was hypothesized that both drugs would inhibit E7-mediated pRb degradation and E7-mediated p21 degradation, resulting in decreased cell cycle progression, immortalization, and proliferation. In addition, we expect that the direct inhibitory action of valproic acid in E7 will target the CKII-mediated phosphorylation pathway necessary for destabilizing p130 and pRb. According to the results of the dynamic simulation, stable interactions exist between every compound. Despite the instability of E7 protein, stability results indicate that both natural chemicals are preferable, with thiazolo outperforming valproic acid.

Key words: anti-cancer, cervical cancer, HPV-16 E7 protein, molecular docking, molecular dynamics simulation.

INTRODUCTION

Human papillomaviruses (HPVs) are a DNA virus family that includes more than 200 different types with a strong preference for squamous epithelium (Tomaić 2016, Bzhalava et al. 2013, Graham 2017). However, they have a similar genetic make-up, various HPVs attack epithelia at different anatomic sites. The anogenital and oral mucosa infections are caused by around 30 different HPV types, which can be further classed as low-risk or high-risk based on the clinical prognosis of the lesions they cause (Münger et al. 2001, Ohlenschläger et al. 2006).

While HPVs that are low-risk generate benign epithelial hyperplasia, HPVs that are high-risk create lesions with a high proclivity to proceed to malignancy. Cervical cancers are commonly linked with high-risk HPV infection in nearly all instances, and two viral proteins, E6 and E7, which are persistently expressed in the transformed phenotype, are essential for developing and maintaining the transformed phenotype, respectively (Shin et al. 2009, Münger et al. 2001). Most research on HPV inhibitors has focused on those oncoproteins due to their well-characterized activities pathways and highly...
differentiate high-risk from low-risk variants (Aarthy & Singh 2021).

The HPV-16 E7 oncoprotein is a 98-amino-acid-long acidic oncoprotein with a calculated molecular weight of around 11 kDa. It possesses strong transforming and carcinogenic capabilities that interfere with normal differentiation and cell cycle control (Valdovinos-Torres et al. 2008, Shin et al. 2009). The E7 protein's three-dimensional structures have been identified, revealing an inherently disordered amino terminus and a novel metal-binding fold in their carboxyl terminus. Conserved regions 1, 2, and 3 (CR1, CR2, and CR3) are composed of two different functional domains at the N-terminus and a single functional domain at the C-terminus (Pang & Thierry 2013, Mok et al. 2018). The CR1 (amino acids 1–18) and CR2 (amino acids 19–42) segments are naturally unfolded with a high degree of flexibility and share structural and functional similarities with the adenovirus E1A protein and the SV40 large T-antigen (Tomaić 2016, Chellappan et al. 1992, García-Alai et al. 2007, Nogueira et al. 2017). The CD1 domain is required for E7 to cause S-phase progression and cellular transformation, whereas the CR2 homology domain comprises the retinoblastoma (pRB) tumor suppressor core-binding site, the LxCxE domain, a casein kinase II phosphorylation site, and its recognition site (Tomaić 2016, Bello-Rios et al. 2021, Knapp et al. 2009, Jones et al. 1997). The CR1 and CR3 have previously been implicated in destabilizing pRB-related proteins p107 and p130 by E7. The C-terminal domain contains the CR3 region, which is structurally similar to the C-terminal domain of E6 oncoprotein and contains two CxxC zinc-binding motifs separated by 29 residues required for zinc-dependent dimerization and protein stabilization (Ohlenschläger et al. 2006, Mok et al. 2018). Additionally, this domain serves as a platform for direct interaction with various tumor suppressor proteins, including p21, p27, and HDAC1. Also, the CR3 region interacts with the C-terminal region of pRB, supporting its degradation (Tomaïc 2016, Todorovic et al. 2011). The full-length E7 protein has a 100-fold greater affinity for pRB than the E7 protein lacking CR1/CR2, indicating a critical function for CR3 in pRb interactions and degradations (Syafitri et al. 2021).

Numerous treatment options targeting the HPV’s main oncoproteins have been developed, including therapeutic vaccines targeting E7 protein, genome editing with antisense oligonucleotides, ribozymes, siRNA, immunotherapy with synthetic E7 DNA sequences, a proteasome inhibitor, and tumor-infiltrating T cells (Syafitri et al. 2021, Tan et al. 2012, Pal & Kundu 2020). However, they are too expensive for the world’s poorest and most vulnerable population, mainly in the middle- and low-income countries (Huh et al. 2017). Despite decades of study, no effective and conclusive therapy for chronic HPV infection has been established yet (Toots et al. 2017). As a mega biodiversity country, Indonesia offers abundant resources that may be developed medically and cost-effectively as an alternative to currently available pharmaceuticals or therapies (Syafitri et al. 2021, Salim & Munadi 2017). We examined asarinin and thiazolo from *Zanthoxylum spp.* and *Myristica fragrans*, respectively, as possible HPV-16 E7 protein inhibitors based on our prior study on HPV-16 E5 and E6 protein inhibitors (data not shown) (Asika et al. 2016, Zhang et al. 2002). There is currently no medicine that precisely targets the E7 protein, thus we utilize a histone deacetylase inhibitor (HDACi), valproic acid, as a control in this *in silico* study. Valproic acid is generally used to treat epilepsy, bipolar disorder, and to avoid migraines, as well as menstrual abnormalities, polycystic ovaries, and hyperandrogenism (Aronson 2015, Andoh
et al. 2020, Feng et al. 2016). Some studies also suggest that it also has anticancer action, notably as an antineoplastic drug identified in 1997 by promoting apoptosis and inhibiting angiogenesis in tumor cells (Cinatl et al. 1997). It suppresses the development of HPV-positive SiHa, CasKi, and HeLa cell lines in vitro, as well as the expression of E7 mRNA. However, its structural effects on the HPV-16 E7 protein remain unknown (Tan et al. 2012, de la Cruz-Hernández et al. 2007, Dai et al. 2017, Feng et al. 2016). This study investigates the potential of asarinin and thiazolo as HPV-16 E7 protein inhibitor candidates using a few molecular dynamic parameters to determine their further potential.

**MATERIALS AND METHODS**

**Data retrieval and pre-docking screening**

The HPV-16 E7 amino acid sequence was obtained from UniProt under the accession number P03129, and the three-dimensional structure of the protein was simulated using the I-TASSER webserver (https://zhanglab.dcmmb.med.umich.edu/I-TASSER/). The 3D model was selected based on its position in C- and TM-score values rankings, among other factors. Using the Ramachandran Plot Server (https://zlab.umassmed.edu/bu/rama/), we evaluated the model’s structure, with the favored percentage coming out to 69.767%. In this study, asarinin (ID 11869417) and thiazolo (ID 1823738) were examined, as well as valproic acid (CID: 3121) as a positive control.

**Molecular docking process**

We used AutoDock Vina integrated within PyRx (https://pyrx.sourceforge.io/) to dock our modeled HPV-16 E7 protein (Hidayatullah et al. 2023). In this study, we look at the whole structure of the target protein, with a molecular coverage area of 36.8165×29.3955×42.2397 Å and center coordinates 51.3889×51.4450×51.5934 Å. The major docking parameters are the molecule’s affinity (measured in kcal/mol), the binding site pocket, and the interaction between the protein and the ligand as our previous study (Widiastuti et al. 2023, Putra et al. 2023).

**Molecular dynamics simulation**

We evaluated all of the protein-compound complexes at 37 °C, 1 atm, pH 7.4, and 0.9 % salinity, which are typical physiological conditions. For 1000 picosecond simulations, the protein and ligand complexes’ structures were generated. A macro program called Md run is used to run a molecular dynamics simulation. The simulation results were examined using the md analyze and md analyzers macro programs through yasara (Hidayatullah et al. 2022).

**Target protein prediction**

Three ligands chosen needed to be validated its protein target, so that the interaction between ligands and protein could be confirmed. To predict the protein target, we utilized SwissTarget webserver (http://www.swisstargetprediction.ch/). The isomeric SMILES of asarinin, thiazolo, and canonical SMILES of valproic acid were used as input (Table I).

**Network analysis and GO prediction**

To predict the interaction of three candidate ligands towards protein target in HPV cases, we performed networking analysis using STITCH database webserver (http://stitch.embl.de/). The webserver automatically generating the Gene Ontology including biological process, molecular function, and KEGG pathway (Heikal et al. 2021).
RESULTS

Before 3D analysis using PyMOL, we align the structure of the E7-ligand complexes before and after the MD simulation with a cycle value parameter of 5.0 and a cut-off value of 2.0 (Fig 1). The alignment results showed that the three complexes deviated around 3-4 Å. As expected, the most significant deviation was experienced by the CR1 and CR2 regions in each complex. Alignment result suggests the E7-thiazolo complex experienced the slightest deviation during the MD simulation process (RMSD: 3.910 Å), while the E7-valproic acid complex experienced the most significant structural deviation during the simulation process (RMSD: 4.942 Å). Deviation in the CR2 region is thought to play a role in causing changes in amino acid residues that interact with the three compounds tested, predominantly shifting more towards the CR3 region. These results are also reflected in the 2D visualization results, which show that less than 55% of pre-MD interactions remain intact after the simulation. Valproic acid only retains its 10% interaction before simulation, whereas asarinin and thiazolo retains some substantial amount of interaction after the simulation, around 40% for asarinin, and 54.5% for thiazolo.

Although the three protein-ligand complexes tested contained less than 55% of the conserved residues, post-MD simulation results revealed that all tested compounds interacted with the same domain as before, with the exception of thiazolo, which interacts with additional domains and critical residues after simulation. While asarinin retains contacts

![Figure 1. The 3D and 2D structure visualization of HPV-16 E7 and (a) asarinin, (b) thiazolo, (c) valproic acid complex. The cyan structures in 3D visualization indicate the protein-ligand complex before MD simulation, while the green structures are after MD simulation. The colored diagram in 2D indicates protein-ligand complex after MD, while the greyscale diagram indicates protein-ligand complex before MD.](image)
with pRb binding domains (Tyr52, Ile76, Arg77), CKII phosphorylation recognition site (Glu35), and p21 binding residues (Ile54), a change in the interacting residues shows when compared to interactions in these three domains prior to the simulation. Valproic acid retains its contact with the pRb binding domain (Tyr52, Arg77) but switches its contact with the CKII phosphorylation recognition sites (Glu35, Asp36, Glu37) to direct interaction with the CKII phosphorylation site (Ser32). On the other hand, thiazolo interacts not only with the p21 binding domain (Ile54, Val55, Leu82), the pRb binding domain (Asn53, Val55), and the CKII phosphorylation recognition site (Glu35, Asp36) but also with an additional critical residue, the HDAC1 binding residue (Leu82).

We also noted that asarinin and thiazolo also remain in contact with the protein’s hydrophobic core. The potential shifting of asarinin causes the compound to interact with more hydrophobic core residues (Ile76, Val69, Ile54). After simulation, the thiazolo interacts with two hydrophobic core residues, namely Ile54 and Leu82. It retains the same amount of contact with hydrophobic core residues as before simulation, but interaction with Ile76 is now with Leu82.

Molecular dynamics simulation for 1000 ps was carried out to predict the stability, interaction pattern, and behavior of each protein-ligand complex with parameters of the physiological condition of the infected person. The energy potential graph shows that the E7-valproic acid complex has the lowest trend value, but it is insignificant compared to the other two complexes (Fig. 2). In addition, the three complexes also show a horizontal trend of graph values throughout the simulation period, around $-3.13\text{e}^{5}$ kJ/mol, proving that the modeled complexes were energetically stable during the MD run.

The RMSD profile (Fig. 3A) of the three control complexes shows a gradual increase in the RMSD value in the first 300 picoseconds and reaches equilibrium. The mean RMSD values of the three complexes were in the range of 3.940-4.66 Å. It is clear that the RMSD graph of the complex E7-thiazolo has a lower value than the other two complexes, as well as fluctuations the minimum during the simulation period. However, all complexes showed the most prolonged equilibration period, confirming the protein’s prolonged stability. The root means square fluctuation (RMSF) was then calculated.
to understand how the mutations affected protein backbone motion.

The RMSF plot (Fig. 3b) shows that the most significant fluctuations occur in the CR1 and CR2 regions of the three protein-ligand complexes. The CR3 region was significantly more stable than CR1 and CR2 over the simulation, universally. Apart from those results, the graph of thiazolo (1.634±0.843 Å) appears to be more stable than asarinin (2.022±1.041 Å) and valproic acid (1.875±0.882 Å). The most significant fluctuations occurred in the region around the LxCxE domain, the CKII phosphorylation site, and its recognition domain, also with the N-terminal of the CR3 region. Those regions are the dominant interaction areas of the three tested compounds before the simulation, predominantly around residues number 30-38 and 52-55 (Table II). However, after simulation, all tested compounds mainly shifted to interact with the CR3 region instead, with better structural stability. These results were also confirmed by the results of the ligand movement simulation (Fig. 3c), which showed extensive ligand movement beyond 5 Å each. Thiazolo offered a stability pattern similar to the previous simulation results and has the most stable ligand movement value among the three compounds tested (5,635±1,049 Å). The most significant ligand movement was experienced by asarinin (7,523±1,339 Å), which experienced significant fluctuations in the first 100 ps of the simulation period and on the timeframes of 750 ps and 950 ps. The three tested ligands appeared to have settled at the most stable binding site after 100 ps simulation with minor fluctuations during the simulation period. The ligand conformation plot (Fig. 3d) shows asarinin has the lowest value of conformational change (1.661±0.312 Å). In contrast to the results of the two previous simulations, which were highly

![Figure 3](image-url)
favored thiazolo, this compound had the highest conformational change value (1.859±0.254 Å). The three compounds tested predominantly had conformational fluctuations around 1.5 to 2 Å.

The 1000 ps SASA simulation revealed that none of the three protein-ligand complexes expanded or shrunk rapidly over the simulation period (Fig. 4a). The E7-thiazolo complex had the flattest and stable trend of SASA values with fluctuations just below 7000 Å² throughout the simulation period lasts. The E7-asarinin and E7-valproic acid complexes experienced two significant changes in two different timeframes. The first fluctuation occurs at 0-300 ps, while the second fluctuation occurs after 800 ps. The E7-asarinin complex has the most drastic fluctuation under 300 ps among all of them. After 800 ps, E7-asarinin and E7-valproic acid start to divert into the different trajectory, relative to flat E7-thiazolo complex. The E7-valproic acid complex tends to shrink after 800 ps. It is reflected with its SASA value, which tends to decrease to 6500 Å² at the end of the simulation. The E7-asarinin complex tends to expand, imaged with the SASA value trend to increase to around 7500 Å² at the end of the simulation period.

The radius of gyration (Rg) values for the HPV-16 E7 protein’s backbone atoms were determined and plotted against simulation time (Fig. 4b). In general, the fluctuations of the three protein-ligand complexes were 13.7-14.6 Å. It is clear that E7-thiazolo complex has the lowest radius value and is stable during the simulation period (13.80±0.14 Å). Another two protein-ligand complexes fluctuate more vigorously, especially the E7-asarinin complex, which jump from 13.5 to 14.6 Å at 100-250 ps timeframe and continue to

Figure 4. (a) SASA plot, (b) Rg plot, (c) intramolecular hydrogen bond plot, and (d) protein-solvent hydrogen bond plot for HPV-16 E7 protein complex with asarinin (red line), thiazolo (green line), and valproic acid (black line) over a 1000 picosecond of simulation.
fluctuate even further to 14.76 Å at the end of the simulation period, around 10% more compared to its initial state. The E7-asarinin complex is also predicted to have an upright trajectory corresponding to its SASA plot. The E7-valproic acid, on the other hand, did not show a plateau graphic pattern showing the stability of the Rg value either and fluctuated around 13.36-14.38 Å. However, it had a lower and more stable mean value (14.02±0.23 Å) compared to the E7-asarinin complex (14.20±0.28 Å).

The intramolecular hydrogen bond plot shows fluctuating values (Fig. 4c). The highest fluctuation was observed in the E7-valproic acid complex, while the E7-asarinin complex and E7-thiazolo complex showed the highest fluctuation, but also a horizontal trajectory even though the graph is not strictly planar. Although the intramolecular hydrogen plot shows relatively significant fluctuation values, it is not apparent in the protein-solvent hydrogen plot (Fig. 4D). The three protein-ligand complexes showed a flat graphic pattern, except for the E7-valproic acid complex, which showed a planar trajectory pattern but tended to decrease even though it was less significant during the 1000 ps simulation.

The output of the DCCM plot revealed that all three complexes exhibited the same pattern of correlated movement (Fig. 5). A series of correlating motions concentrated on the CR3 region, specifically in the β₁ and β₂ structures, as well as an α-helix immediately behind them. The E7-valproic acid complex appears to have far more correlated mobility than the other two complexes. Although the movement intensities of the E7-asarinin and E7-thiazolo complexes were almost comparable, the E7-thiazolo complex exhibited a relatively lower movement intensity, indicated by lower yellow intensity on the plot.

In order to validate the function and cascade pathway of three candidate ligands, we performed protein target prediction (Supplementary Material - Figure S1), continued with networking analysis (Fig. S2 and S3). The protein target of asarinin shows that MCL1 (Induced myeloid leukemia cell differentiation protein Mcl-1) has the highest probability score as target (0.78). While target protein of thiazolo has similar probability score around 0.1, including ENPP1, ADORA1, ADORA2A, ADORA2B, PTPsigma, DHODH, CRHR1, ELANE, GPR39, CCNB3. Valproic target protein probability score varied

![Figure 5. Dynamic cross-correlation matrix for each protein-ligand complex. Yellow represents correlated motions, and blue represents anticorrelated motions.](image-url)
between 0.07 to 0.14. Top three highest target protein score are PPARD, FFAR1, and FABP2.

Target proteins of asarinin included in various pathway, mainly in stress regulation, binding with protein and transcription factor, and playing roles in cancer pathway, renal and colorectal carcinoma, hepatitis B and HIF signaling pathway. While GO analysis of thiazolo’s target protein mainly acts in the phosphorylation activity, signaling activity, and included in the cell cycle, viral and cancer pathway (Table III and IV). The terminology did not show literal name as HIV inhibitor pathway.

The network analysis did not show any interaction between compounds and protein targets. This probably because there is no in vivo/in vitro experiment proven the protein targets of asarinin and thiazolo directly (Fig. S2 and S3).

**DISCUSSION**

Molecular docking and dynamics simulations indicate that asarinin, thiazolo, and valproic acid are primarily localized in the C-terminal of CR2 and CR3 regions. It is assumed that these two areas are more stable and less flexible than the CR1 and N-terminal CR2 regions (Chellappan et al. 1992, Nogueira et al. 2017, Calçada et al. 2013). The very flexible and unstructured nature of the CR1-CR2 region was further demonstrated by the comparatively high pre-and post-MD superimposed root mean square deviation alignment values (>2 Å), also further by the RMSF plot. The flexibility of the two N-terminal domains is also suggested to explain why valproic acid loses 90% of its contacts with E7 protein residues following simulation, whereas the other two compounds lose more than 50% of their initial associations with E7 protein residues. It suggests that the main reason for supporting the main important domain for E7 protein oncogenic activity is that the LxCxE motif in CR2 is available for interaction with pRb without requiring extensive structural reorganization (Ohlenschläger et al. 2006).

The three simulated compounds interacted with two key residue complexes, the pRB binding site and the CKII recognition domain, in a very similar manner. Although the major contact between E7 protein and pRB occurs in the LxCxE domain of the CR2 region, an interaction domain with pRB exists in the CR3 area as well. The interaction in the CR3 area exhibits a lesser affinity when compared to the interaction in the LxCxE domain. However, in the absence of interaction with CR3, the E7 protein’s affinity for pRB can diminish significantly. The three compounds interact primarily with the pRB binding site in the CR3 region through Tyr52, Asn53, Val55, Ile76, and Arg77. All contacts at the binding site are mediated by hydrophobic interactions, which are more robust than hydrogen bonds in the context of weak molecular force. Thus, the interaction between asarinin and thiazolo, and valproic acid with protein E7 is relatively stable and is predicted to be capable of decreasing its affinity for pRb, resulting in a slower rate of degradation of the pRb-E2F complex, which is inadequate to offset the effects of pRb-induced cell cycle arrest (Nogueira et al. 2017, Todorovic et al. 2011).

According to the simulation results, both natural compounds exhibited sustained interactions with the CKII recognition domain (aa 33–37). The domain is a five-residue expansion of the C-terminal CKII phosphorylation site. Disruption of the recognition domain disrupts the phosphorylation of pRb at Ser31 and Se32, resulting in a reduction in the affinity of E7 protein for the RbAB domain on pRb protein. This phosphorylation step is critical in the method by which the E7 protein regulates cycle cell control since phosphorylated pRb by CKII interacts...
with E2F transcription factors, repressing their activity and reducing DNA synthesis (Chemes et al. 2010, Basukala et al. 2019). Prior to that, the CKII required recognition of the specific location (EEEDE) in order to interact with E7 and phosphorylate pRB. Inhibition at this location impairs the protein’s phosphorylation ability (Phelps et al. 1992). However, the importance of CKII-mediated E7 phosphorylation on pRB binding and instability has been disputable (Chemes et al. 2010, Genovese et al. 2008, Firzlaff et al. 1991).

Additionally, valproic acid interacts with one of the CKII phosphorylation site’s serine residues (Ser32). The interaction, which is mediated by hydrophobic interaction and a moderate strength hydrogen bond, is thought to be relatively stable and thus would directly reduce E7’s affinity for CKII, impairing its phosphorylation capabilities even further, resulting in even less pRB being phosphorylated, impairing the cells’ immortality and replication (Jones et al. 1997, Chemes et al. 2010, Basukala et al. 2019, Genovese et al. 2008, Phillips & Vousden 1997). Additionally, interventions on this group of domains disrupt the CKII-mediated phosphorylation process necessary for destabilizing Retinoblastoma-like protein 2 (RBL-2) or p130, which is essential for promoting infected cells into the S phase via E2F-mediated transactivation (Genovese et al. 2008, Cobrinik 2005, McLaughlin-Drubin & Münger 2009).

Within the CR3 region, asarinin and thiazolo hypothesized to interfere with dimerization and stabilization mechanism of the E7 homodimer complex due to their interaction with the hydrophobic core of the CR3 region. They are critical in E7 protein dimer structure stabilization, particularly in the intermolecular contacts between the two-stranded antiparallel β-sheets on each monomer (Ohlenschläger et al. 2006, Todorovic et al. 2011, Liu et al. 2006). Meanwhile, thiazolo is also predicted to interact with a key residue of HDAC1 binding site in the region; Leu82, which overlapped with E7’s hydrophobic core residue. It is hypothesized that disrupting Leu82 and Leu83 would abolish HDAC1’s association with E7 protein and its capability to transform because altering chromatin structure occurs via deacetylation of histones, which play a critical role in HPV E7-associated transcriptional control and can extend the life of infected cells (Brehm et al. 1999, Lourenço de Freitas et al. 2021, Longworth & Laimins 2004, Helt & Galloway 2001).

Meanwhile, the p21 binding domain in CR3 overlaps with the low-affinity pRb binding region, likely since a portion of p21’s C-terminus shares a sequence with the pRb’s C-domain (Ohlenschläger et al. 2006, Liu et al. 2006). Thiazolo interacts with more residues (Ile54, Val55, Leu82) than asarinin, which may affect their inhibitory capacity, which is favored for thiazolo. Both of them are predicted to reverse p21-mediated inhibition of cyclin E–CDK2 and cyclin-A–CDK2 kinase activity and contribute to HPV E7’s ability to reestablish and sustain CDK2 activity in differentiating keratinocytes, thereby allowing for viral replication by trying to interfere with CDK2 that could significantly affect Rb-dependent E2F transcription regulation, as well as pRb phosphorylation. It is hypothesized that disrupting the E7-p21 interaction would impair the E7 protein’s capacity to stimulate DNA synthesis in differentiated cells (Shin et al. 2009, Jones et al. 1997, Parveen et al. 2016, Boshoff & Weiss 2002, Lehoux et al. 2009).

The potential energy in the three complexes was analyzed, and it was discovered that the three complexes remained stable over the simulation time. It means that the three complexes’ electrostatic and van der Waals interactions remained constant during simulation, suggesting that no anomalies
occurred during the simulation process (Bavi et al. 2016, Albaugh et al. 2016). When compared to the other two complexes, the RMSD analysis suggests that the E7-thiazolo complex has the most stable backbone (Aier et al. 2016). During the simulation time, however, all three displayed a similar stabilization pattern, showing that they did not suffer considerable deformation after reaching their equilibrium state, despite their value exceeding 2 Å, owing to the instability of CR1 and CR2, as corroborated by the RMSF plot (Aier et al. 2016, Sivaramakrishnan et al. 2020, Ohlenschläger et al. 2006, Tomač 2016). The RMSF plot reveals that all compounds’ initial binding site residues are highly flexible, contributing to extensive ligand movement and conformational changes. Each of those compounds changes conformation and location in response to the dynamics of its binding site during the simulation process (Nogueira et al. 2017, Calçada et al. 2013, Zhu et al. 2017, p.1).

Protein integrity indicators such as SASA, Rg values and hydrogen bond diagrams reveal that the integrity of all protein-ligand complexes is rather constant over the simulation period and exhibits no indications of denaturation (Zhang & Lazim 2017). These factors favor E7-thiazolo yet again. The SASA plot reveals that none of the complexes change dramatically in a short amount of time, but E7-asarinin has an upward trajectory, indicating that it is about to expand. On the other hand, E7-valproic acid displays a downward trajectory, indicating that it has reduced in size relative to its natural structure (Zhang & Lazim 2017, Kamaraj & Purohit 2013, Sneha & George Priya Doss 2016). The Rg analysis, which assesses the rigidity of the protein structure, shows that the thiazolo, and valproic acid complexes are stiffer than the asarinin complex because they have a lower Rg value than asarinin (Dash et al. 2019, Sneha & George Priya Doss 2016). Intramolecular hydrogen bond analysis revealed relatively extensive movement.

In contrast, protein-solvent hydrogen bond analysis revealed a relatively stable pattern, presumably due to critical factors such as E7’s size, CR1, and CR2 that naturally unfolded, and they moved continuously over the simulation time to rearrange themselves to achieve equilibrium (Liu et al. 2006, Ohlenschläger et al. 2006, Petukhov et al. 2004, Kuhn et al. 2010). However, they still retain their hydrophobic core, retaining the structure and staying folded (Mallamace et al. 2018, Fitzpatrick et al. 2011). Because all secondary structures are located within the CR3 region, the DCCM analysis demonstrates that all associated motions are centered inside this region. Given that both the asarinin and thiazolo complexes exhibit a similar degree of correlated movement, indicating that they are less elastic than the valproic acid complex, it is reasonable to presume that both complexes are more stable than valproic acid (Valente et al. 2020, Dash et al. 2019).

The protein targets prediction and networking analysis successfully generating some terminology related to cancer pathway and phosphorylation activity. Even though only one terminology related to viral carcinogenic found in thiazolo’s gene ontology, this could be the potential of carcinogenesis development initiated by viral infection. Each virus promotes carcinogenesis in its unique way. The majority of DNA oncogenic viruses encode oncogenes that change infected cells, with p53 and pRB being the most common targets. In addition, for certain viruses, including HBV and HPV, viral DNA integration into the human genome can play a key role in tumor growth. Because viral integration necessitates the breakage of both viral and host DNA, the rate of integration is thought to be correlated with DNA damage levels. Endogenous and exogenous causes, such
as virus-induced inflammation or co-infections with other agents, environmental agents, and other factors, can cause DNA damage. Cancer usually takes years or decades to develop after an infection (Chen et al. 2014). Cervical cancer, penile cancers, and several other anogenital and head and neck cancers are linked to high-risk human papillomaviruses (HPV) 16 and 18 (some other -HPV variants are also carcinogenic) (Boshart et al. 1984, Dürst et al. 1983).

CONCLUSIONS
The HPV-16 E7 protein is one of the most well-known oncoproteins in cervical cancer cases directly involved in cell cycle progression, proliferation, and immortalization through its interaction with pRb and the pRb-related p107 and p130, CKI such as p21, and histone deacetylase. Through molecular docking and dynamic simulation, we directly target it with asarinin and thiazolo. We predict both compounds could inhibit E7-mediated pRb degradation as well as E7-mediated p21 degradation, which leads to cell cycle progression, immortalization, and proliferation. Also, we predict that the direct inhibition mechanism of valproic acid in E7 is to target the CKII-mediated phosphorylation process necessary for destabilizing p130 along with pRb. Dynamic simulation results suggest that all compounds had stable interactions. Despite the unstable nature of E7 protein, the stability results favor both natural compounds, particularly thiazolo, over valproic acid.

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SUPPLEMENTARY MATERIAL

Figure S1-S3.

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Author contributions

A.H.; W.E.P.; S.; and M.R. planned and designed the research. A.H. performed molecular docking. W.E.P. performed molecular dynamics simulation. G.W.P. performed target protein and network analysis. W.E.P.; S.; M.R.; D.W.; M.F.H. analyzed the data. A.H. and W.E.P. wrote the manuscript. All authors revised the manuscript.