

Theoretical Evidence for the Antibacterial and Antioxidant Potentials of a New Molybdenum Tricarbonyl Complex

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Abstract

The development of low energy light-activated therapeutics with precise targets is currently receiving pronounced attention. Here we report the evaluation of a novel mer-tricarbonylmolybdenum(0) complex with benzyl isonicotinoyl hydrazone (BIH) as ancillary ligands. The complex is a photoactive carbon monoxide-releasing molecule (PhotoCORM) with dual action. Spectroscopic characterization (IR, NMR) and the elemental analysis results confirmed a meridional octahedral geometry tricarbonyl complex with a para substitution pattern for the coordinating pyridinic nitrogen atom of the ancillary ligand. FTIR spectral analysis strongly suggested that the para substitution mode of the ligand attenuated metal-to-ligand π -backbonding, resulting in the weakening of the Mo-C bond. Time-dependent density functional theory (TD-DFT) calculations showed that a highly allowed Metal-to-Ligand Charge Transfer (MLCT) in the visible light region ($\lambda_{max} = 587.45$ nm) initiated the photolysis process. Experimental evidence supported that upon irradiation, the complex undergoes dissociation at a rapid rate ($k = 9.7 \times 10^{-2} \text{ s}^{-1}$). Released CO is successfully delivered to biological heme sites as confirmed by carbonmonoxy myoglobin (MbCO) formation. *In silico* molecular docking simulation with SwissDock program demonstrated that possible release of hydrazone ancillary ligand may effectively target the active site of keap1 (PDB: 4L7B) and those of essential bacterial proteins; LasR (PDB: 3IX3), MRSA PBP2a (PDB:5M18) and *E. coli* DNA Gyrase (PDB: 6F86). *In vitro* antibacterial assay carried out to validate the theoretical broad-spectrum activity of the ligand, and the CO poisoning of the heme indicates significant growth inhibition against *S aureus*, *E. coli*, and *P. aeruginosa* in a dose-dependent manner. These results highly suggest this novel photoCORM as a promising synergistic antimicrobial agent.

Keywords: photoCORM, TD-DFT, molybdenum tricarbonyl, molecular docking, CORM antibacterial

1. Introduction

Substituted transition metal carbonyls are recognized as the best class of carbon monoxide releasing molecules (CORMs). With carefully selected metal and biologically active ancillary ligands they display better antibacterial activity compared to the homonuclear starting complex

[1]. We have previously reported the antibacterial activities of tungsten and molybdenum tricarbonyls designed with hydrazones as ancillary ligand [2,3]. They showed remarkable activity against Gram-positive and Gram-negative pathogens. Mechanistic studies implicate CO release by CORMs in physiological conditions as a trigger for antibacterial action. Such CO released by

CORMs is suspected to impair bacterial respiratory pathway as the major cause of antibacterial action [4].

Several CORMs have been reported to have impressive antibacterial activities [5]. Although the mechanisms of CORMs' antibacterial action are not entirely understood, it is thought to include disruption of bacterial membranes, possibly due to hydroxy radical production, interference with metal ion uptake in bacteria, and CO binding to respiratory cytochromes, which may hinder bacterial respiration [1]. Major challenges in achieving useful antibacterial CORMs include tissue-specific CO delivery and toxicity, as CORMs can be toxic at higher doses [6]. There is therefore a need to devise strategies for optimizing carbon monoxide release from CORMs. This can be achieved by a careful selection of tunable ancillary ligands, metal centre of desirable redox potential, stability, and biocompatibility.

CORMs have elicited burgeoning scientific interest and some debates. A recent investigation reported that water-soluble CORM-2, which is a potential antibacterial agent, kills bacteria both in culture and animal models but is benign towards mammalian cells [7]. According to the researchers, CORM-2 releases less than 0.1 M CO per mol, which is negligible, and this challenges the prevailing notion that CO is responsible for the antibacterial activity of the CORM. Ruthenium accumulation is fingered for its antibacterial activity and not CO release.

Theoretical computational chemistry promises a pivotal role in advancing our understanding of the properties, bioactivity, and mechanism of CORMs. However, this application is currently underexplored. We can gain insight into CORMs' CO release kinetics and estimate CO liberation rates with knowledge of the thermodynamics and reaction pathway parameters [8,9]. Quantum mechanical optimization and frequency calculations can be very valuable in the design of CORMs as a computational tool in predicting stability, reactivity, CO release, and binding affinities. Density functional theory (DFT) has been used to elucidate metal-CO bonding, giving insights into metal carbonyl complexes [10]. Computation models and docking studies with pathogens are reported to provide information regarding toxicity and antibacterial targets. DFT calculations performed using the B3LYP/LANL2DZ basis set were used to explore the CO release capacity of some manganese-containing CORMs [11,12]. The calculations predicted the Mn-CO bond dissociation energies, providing insight into the amount of CO released by the CORMs. The natural bond orbital (NBO) analysis revealed the donor-acceptor interactions and bond stabilization energies of Mn-CO

within the CORMs studied. Impressively, the study analyzed the nucleophilic attack of putative ligands at the metal vacant site using ligand-exchange reaction energies. Calculated second dissociation energies provided insights into the observed stoichiometry [8,13]. Similar DFT studies of some group 7 CORMs bearing pyridine-based moieties as ancillary ligand compared the electronic properties of manganese and rhenium tricarbonyl complexes in terms of their carbon monoxide (CO) releasing tendency. Calculated absorbance, energy gap, and metal-to-ligand charge transfer (MLCT) percentage were employed in characterizing the tendency of the tricarbonyl complexes in labilization of CO [14].

Both experimental and theoretical analyses have been used to study Mn PhotoCORMs; Mn(I) tricarbonyl complexes containing the di(2-picoly)amine and similar units with various electron-withdrawing ligands. DFT calculations and TD-DFT reveal their CO-releasing potential [12,13,15]. Electronic transition calculations reveal a robust MLCT, which weakens the metal-CO back bonding and facilitates rapid CO release. Similarly, TD-DFT calculations using both BP86 and B3LYP functionals were employed to study electronic properties, molecular orbitals, and excitation energies. These reveal information about the electronic structure and CO-releasing behavior of the complexes [16]. Molecular docking studies have proved to be useful in the investigation of CO-releasing coumarin derivatives as an antitumor agent. However, photoCORM research is currently challenged with issues of light activation wavelength and tissue penetration, aqueous stability, toxicity of central metal atom, targeted delivery etc.

In the current study, we have employed a novel approach of using computational chemistry to get theoretical insights into the structure, reactivity, CO release mechanism, antioxidant, and antibacterial action of a tricarbonyl molybdenum(0) containing benzylisonicotinoyl hydrazone as an ancillary ligand. This can assist in the elucidation of the exact mechanism of antibacterial action. The choice of the Mo(0) tricarbonyl, a d6 species for this study, is intended to address issues of biocompatibility, favourable light activation and kinetic control. The tricarbonyl core can improve CO release lability compared to lower di- and mono-carbonyls. This scaffold if conjugated to an extended π -conjugating system such as the isonicotinoyl hydrazone ancillary ligand will likely improve light activation and CO lability.

2. Materials and Method

2.1. Materials

Molybdenum hexacarbonyl, trimethylamine N-oxide dihydrate (TMNO), isonicotinic hydrazide (INH), benzaldehyde, tetrahydrofuran (THF), acetonitrile, and dichloromethane were purchased from Sigma Aldrich and used without further purification. All other reagents used in this work were all-purpose laboratory reagents. All reactions involving metal carbonyls were performed under pure dry nitrogen gas using manipulated Schlenk techniques. Solvents were of reagent grade and purified according to standard methods.

2.2. Preparation of (E)-N'-benzylideneisonicotinoyl hydrazide (BIH)

The ligand synthesis follows the protocol method reported for the condensation reaction between hydrazides and aldehydes to yield asymmetric Schiff base hydrazone [3]. Isonicotinic acid hydrazide (1.827 g; 0.013 mol) dissolved in ethanol in a round-bottom flask was added benzaldehyde (1.413 g; 0.013 mol) with constant stirring. The mixture was refluxed for 4 h, and the product was collected in a beaker, which crystallized immediately on cooling. The white solid obtained by filtration was recrystallized from ethanol and dried over fused calcium chloride in a desiccator and then weighed (Yield = 73%).

2.3. Preparation of [Mo(CO)₃(BIH)₃]

In a one-pot synthesis, a mixture of Mo(CO)₆ (0.396 g; 1.43 mmol), BIH (1.014 g; 4.53 mmol) previously dissolved in 10 mL of methanol and trimethylamine N-oxide dihydrate (0.339 g; 3.05 mmol) in 25 mL of tetrahydrofuran was stirred under dinitrogen at ambient temperature. An initial yellow solution changed to orange and finally to a reddish-brown suspension. The reaction was allowed to stand for 36 h, and the mother liquor was reduced to a small volume under reduced pressure. The reddish-brown solid was isolated by vacuum filtration and washed with a hexane-dichloromethane mixture and dried over silica gel in a desiccator [17].

2.4. Characterizations

Infrared and Raman spectra were obtained using a SHIMADZU Fourier transform infrared spectrometer using the KBr disc pellets technique and ProRaman-L-785B Raman analyzer, respectively. NMR spectra were recorded using a Bruker NMR-400 MHz spectrometer. UV-1800 Series spectrophotometer configured with a slit width of 1.0 nm and a light source change wavelength of 560 nm was used to record absorbance spectra data.

2.5. Computational Studies

The 2D guess structures of the compounds were drawn in ChemDraw and uploaded to the GaussView16 interface. The optimization and frequency calculations of the clean structures were carried out using the DFT, TD-DFT methods, and B3LYP/6-311+G (d,p) model chemistry in Gaussian09 [18]. The online programs SwissADME [19] and pkCSM [20] were used for the ADMET profiling of the ligands. The molecular docking studies to examine the interactions between the organic ligands against LasR quorum-sensing receptor of *P. aeruginosa* (PDB:3IX3), PBP2a transpeptidase of MRSA (PDB: 5M18), and the DNA Gyrase B subunit of *E. Coli* (PDB: 6F86) [21] and Keap1 4L7B [22] were carried out using the SwissDock program [23], [24]. The 3D structure of the *P. aeruginosa* (PDB:3IX3), MRSA (PDB: 5M18), PDB: 6F86, and PDB: 4L7B were downloaded from the RCSB Protein Databank. The docking results were analyzed using ChimeraX [25,26] and BIOVIA Discovery Studio Analyzer software [27].

2.6. Kinetics of CO Release by Metal Carbonyl Complexes

The substituted molybdenum tricarbonyl (0.005 g) placed in a quartz cuvette, was added to 1.00 mL DMSO solution. The mixture was shaken to ensure complete dissolution of the metal carbonyl, and the absorbance was immediately recorded to determine the λ_{max} . Fresh solutions were then prepared, and absorbance was recorded at the λ_{max} observed for the complex to determine the rates of CO release. Graphical plots of absorbance versus time were obtained. The apparent rate of CO release was calculated from the $\ln[C]$ versus time plot for the substituted tricarbonyl complex.

2.7. CO Delivery to Myoglobin in Solution [28]

Equine heart myoglobin (53 μ M) was dissolved in phosphate-buffered saline (pH 7.4), constituted by dissolving one buffer tablet in 200 mL of deionized water. The solution was reduced by adding sodium dithionite. The concentration of deoxy-myoglobin (deoxy-Mb) generated was calculated to determine the concentration of tricarbonyl added. The initial absorbance was recorded immediately on adding the tricarbonyl complex. For Carbonmonoxy myoglobin (Mb-CO) formation, the cuvette was irradiated using a monochromatic LED light source with maximum centered at 560 nm. The light intensity was maintained at 20 mW/cm² and the spectra was monitored over time at regular intervals.

2.8. Antibacterial Studies of Substituted Molybdenum Tricarbonyl carbonyl

The ^1H NMR spectroscopy perfectly corroborates the FTIR result; the tricarbonyl complex displayed a relatively symmetrical splitting pattern in the aromatic region (7.0–8.8 ppm). This non-clustered splitting is characteristic of a 1,4-disubstituted (para)pyridine ring displaying two

the amide carbonyl of the hydrazone ligand is in highly nucleophilic regions. While in the electrophilic region is centred over the aromatic protons and the N-H of the ligand. This charge polarization suggests the tricarbonyl possesses adequate physiological solubility.

Table 2. Theoretical and experimental values of the percentage of C, H, and N in the complexes.

Compounds	Molecular mass	C		H		N	
		Calc.	Found	Calc.	Found	Calc	Found
$\text{Mo}(\text{BIH})_3(\text{CO})_3$ ($\text{C}_{42}\text{H}_{33}\text{N}_9\text{O}_6\text{Mo}$)	855.73	58.95	58.46	3.89	4.01	14.73	14.40

distinct doublets for the four pyridine protons. The CHN elemental analysis (Table 2) and the ^1H NMR and vibrational spectroscopic data all indicated a meridional tricarbonyl complex with the proposed structure (Fig. 2).

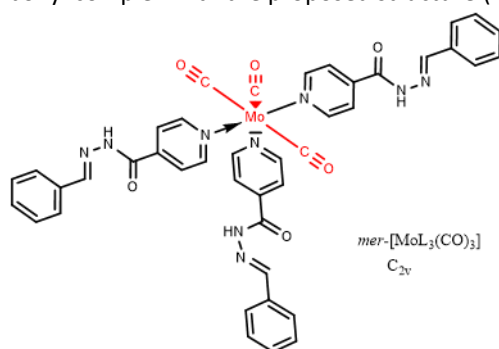


Figure 2. The proposed structure of the tricarbonyl complex.

The geometry and electronic distribution in the molecule, according to the spectral analysis, affect the strength of the Mo-C bond. The rate of CO dissociation in metal carbonyls is largely determined by this M-C bond. According to the FTIR analysis, this para-substituted hydrazone is a weaker electron donor due to the resonance effect of the para electron-withdrawing substituent. This depletes electron density, leading to weaker π -backbonding ($\text{Mo} \rightarrow \text{CO}$). This leaves the Mo-C inherently weaker and more labile, and the thermal or photochemical cleavage required to release the CO will occur more readily and at a faster rate [20].

3.2. Optimized Structure, Global Reactivity Parameters, and Molecular Electrostatic Potential (MEP) Map

The MEP map and global reactivity parameters calculated from the optimized ground state geometry can help in the understanding of the chemical behaviour and molecular interaction profile of the studied tricarbonyl. The MEP map (Fig. 3) revealed that the oxygen atom of the CO coordinated to the Mo atom, and

Table 3. Global reactivity Parameter of mer-Mo (CO) $_3$ (BIH) $_3$.

	HOMO	LUMO	ΔE	μ	η	ω
$\text{Mo}(\text{BIH})_3(\text{CO})_3$	-4.4278	-2.1894	2.2384	-3.3086	1.1192	4.890

The chemical hardness and chemical potential calculated from the HOMO-LUMO values (Table 3) indicate a moderately soft and highly polarizable system. This electronic softness enhances the large instantaneous shift in electron density required during the MLCT to trigger CO release. The electrophilicity index of 4.89 eV suggests a strong propensity for the carbonyl to act as an electron acceptor during initial biological interaction before photolysis.

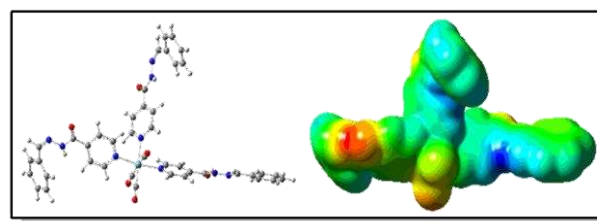


Figure 3. Optimized structure and Molecular Electrostatic Potential Map of the substituted tricarbonyl complex.

3.3. Frontier Molecular Orbital (FMO) Analysis

The spatial distribution and energy of molecular orbitals (Fig. 4) can also provide useful insight into the application of the studied tricarbonyl as a photoactivatable CORM. The calculated energy gap of 2.239 eV is narrow and corresponds to an electronic transition in the visible region ($\lambda_{\text{max}} = 554 \text{ nm}$), which is suitable for biological applications and can allow for deeper tissue penetration with minimal phototoxicity. This is in substantial agreement with the TD-DFT findings, further suggesting the carbonyl's preference to target visible light excitation. The spatial distribution of the frontier orbitals provides mechanistic information regarding photo-induced CO release. The HOMO electron density is heavily localized in the mer-[Mo(CO) $_3$] moiety, revealing the $d\pi-\pi^*$ metal-to-ligand backbonding that stabilizes the tricarbonyl. But the LUMO is localized

almost exclusively on the extended conjugated π -system of the isoniazid hydrazone ancillary ligand in the para position. These electronic events accompanying the lowest-energy excitation constitute a pronounced Metal-to-Ligand Charge Transfer (MLCT), suggesting that upon absorption of a photon by the molecule, electron density is acutely depleted from the Mo centre and transferred to the peripheral hydrazone ligand. The metal core undergoing a transient oxidation leads to a weakening of the M-CO π -backbonding, which kinetically labilizes the Mo-C bond, resulting in controlled release of CO.

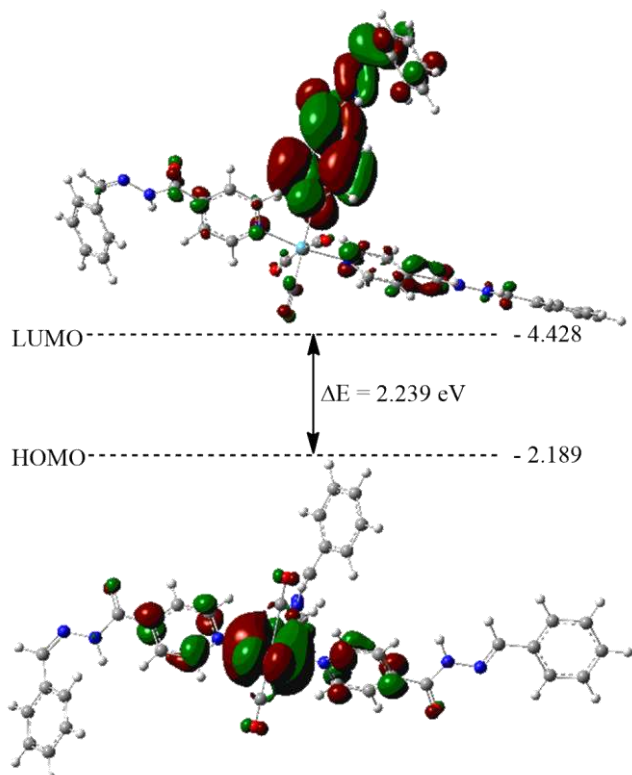


Figure 4. Frontier Molecular Orbitals displaying electron distribution and energy change.

3.4. Time-Dependent Density Functional Theory (TD-DFT) and Electronic Transitions.

The TD-DFT calculations can provide information about electronic transitions, which can be used for photochemical characterization. The theoretical UV-vis spectrum reveals a dominant, highly intense absorption band located within the visible region (Fig. 5). From the theoretical spectroscopic data (Table 4), the transition which occurred at 587.45 with an excitation energy of 2.1106 eV defines the dominant band as the primary absorption maximum driven mainly by the excited state 5. This state is a highly allowed and statistically probable electronic transition state with its high oscillator strength of 0.5157 release corroborate the activation mechanism.

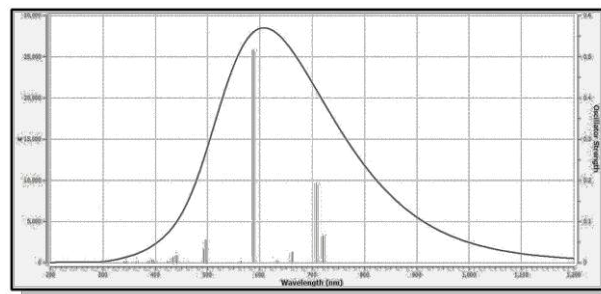


Figure 5. Calculated UV-Visible spectrum of $\text{Mo}(\text{BIH})_3(\text{CO})_3$.

The MO contributions for this excited state revealed that the strong absorption is characterized mainly by the promotion of electronic transition from HOMO -1 (MO 204) to LUMO+2 (MO 208) (Fig. 6) with a dominant expansion coefficient of 0.63401 (80.39 %). The promotion of this election triggers the metal-to-ligand charge transfer required to disrupt the $\text{M} \rightarrow \text{CO}$ backbonding. This strong absorption within the visible light region (yellow-orange light) computationally supports the viability of this substituted tricarbonyl as a PhotoCORM.

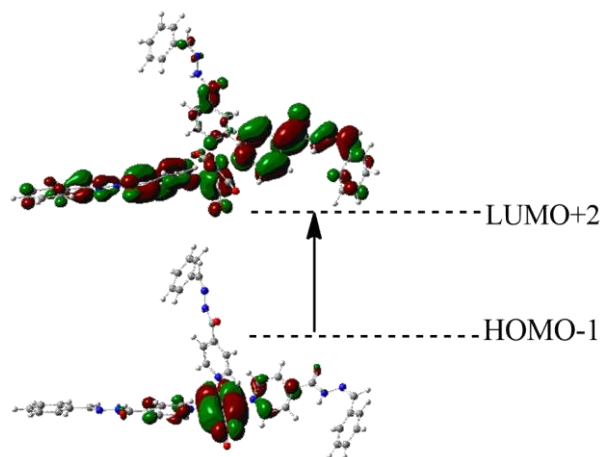


Figure 6. MOs showing the major electronic transition from the Mo center to the ancillary ligands, which give rise to M $\rightarrow$ L charge transfer transitions.

Its low visible light activation potentially eliminates possible phototoxicity and limits tissue penetrations associated with high energy irradiation [28-30]. This absorption led to promotion of an electron from MO204 localized on Mo to MO208 located on the extended π -system of the isonicotinic hydrazone ligand. The migration of this electron density from metal to the ancillary ligand depletes the electron used by Mo for π -backbonding with CO. The Mo-C is thereby weakened facilitating CO release. The dominant peak emerging at 570 nm in the myoglobin assay which is a proof of CO. Additional lower energy transition appearing at 707.80

Table 3. Theoretically calculated energy levels of the lowest-lying transitions of the tricarbonyl.

State	Assignment		Coefficient	Energy (eV)	Wavelength (nm)	Oscillator strength
	From	To				
S0 → S1	HOMO	LUMO	0.6859 (94.09 %)	1.7209	720.48	0.0645
	HOMO	LUMO+2	-0.1399 (3.91 %)			
	HOMO-1	LUMO+2	-0.1868 (6.98 %)			
S0 → S2	HOMO	LUMO+1	0.6579 (86.57 %)	1.7517	707.80	0.1936
	HOMO	LUMO+2	0.1197 (2.87 %)			
	HOMO-1	LUMO	-0.1619 (5.24 %)			
S0 → S5	HOMO-1	LUMO+1	-0.1306 (3.41 %)	2.1106	587.45	0.5157
	HOMO-1	LUMO+2	0.6340 (80.39 %)			
	HOMO	LUMO+1	0.2049 (8.40 %)			

nm further extends the therapeutic optical window for the application of this compound in biological tissues by enhancing its profile for targeted phototherapy. This secondary low energy absorption is an allowed transition, primarily composed of excited State 2 ($E = 1.7517$ eV, $f = 0.1936$).

3.5. The CO release kinetics and delivery to deoxymyoglobin

The quantitative validation of the CO release profile of the tricarbonyl in solution was carried out via time-dependent UV-Vis spectrophotometry. A standard calibration plot of concentration against absorbance was used. Graphical plot of $\ln[\text{Mo}(\text{BIH})_3\text{CO}]_3$ versus time yielded a linear relationship indicating CO release strictly follows first-order kinetics. By fitting of the exponentials in the first order rate equation, the apparent CO release rate (k_{CO}) was determined to be $9.7 \times 10^{-2} \text{ s}^{-1}$. To confirm the delivery of CO to myoglobin, the introduction of the freshly prepared solution of the carbonyl complex to the standard deoxymyoglobin assay at physiological pH (7.4), changed the initial absorption maximum at 405 nm, forming a new dominant peak at 570 nm. This shift serves as a diagnostic tool to prove the complete conversion of deoxymyoglobin to carbonmonoxy myoglobin, validating the capacity and therapeutic potential of the tricarbonyl complex to deliver CO directly to the heme centre.

3.6. Molecular Docking Studies

The CO release by CORM in biological systems plays well established roles. The CO release events however give rise to the generation of secondary species. These species and their actions play key roles in the successful application of the CORM. MLCT driven CO release triggers a dissociation process yielding coordinatively unsaturated metal fragments and free ligand derivatives. The careful choice of isonicotinoyl hydrazone scaffold as an ancillary ligand here is because the light-triggered release can lead to the free ligand permeating the intracellular space to interact with certain protein

targets. Such interactions can enhance antibacterial, anticancer and antioxidant activities. Therefore, in silico molecular docking simulation of the hydrazone is employed to evaluate its action against selected bacterial targets and Kelch domain of Keap1 (PDB: 4L7B) for antioxidant response.

3.6.1. Antibacterial

Molecular docking of the hydrazone scaffold was performed against three high-priority targets, the same as those used in experimental evaluation. The bacterial targets LasR quorum-sensing receptor of *Pseudomonas aeruginosa* (PDB:3IX3), PBP2a transpeptidase of Methicillin-Resistant *S. aureus* (PDB: SM18), and the DNA Gyrase B subunit of *E. coli* (PDB: 6F86) were used. The docking protocol was first validated by redocking the co-crystallized standard ligand inhibitors in the targets. The top scorer redocked ligand pose was superimposed with the proteins' co-crystallized ligands respectively. The matching of the poses showed good RMSD score of 0.95 Å, 0.54 Å, 0.99 Å, and 1.01 Å for LasR, PBP2a, DNA Gyrase and Keap1 respectively. The docking poses (Fig. 7) showed that the hydrazone remarkably conforms to the distinct active sites of both the Gram-positive and Gram-negative targets. The bonding affinity is largely driven by hydrogen bonding via the pyridine N and the hydrazone linker

3.6.2. Antioxidant

Since CO is recognised to upregulate cellular antioxidant responses, it is necessary to investigate if the likely liberation of ancillary hydrazone during CO release could synergistically amplify the process. The ligand was docked against the Kelch domain of Keap1 (PDB: 4L7B), which is the primary negative regulator of the Nrf2 oxidant pathway. The result (Fig. 8) revealed the hydrazone ligand formed two stable hydrogen bonding interaction through the pyridine N with the highly conserved ARG 483, and second through the amide carbonyl oxygen with ARG 415 (2.031 Å). The hydrazone

effectively fits into the shallow regulatory pocket, with a binding score of -7.068, demonstrating its likely ability to sterically inhibit the sequestration of Keap1-Nrf2 process, and potentially acting in direct synergy with the cytoprotective effect of the liberated CO.

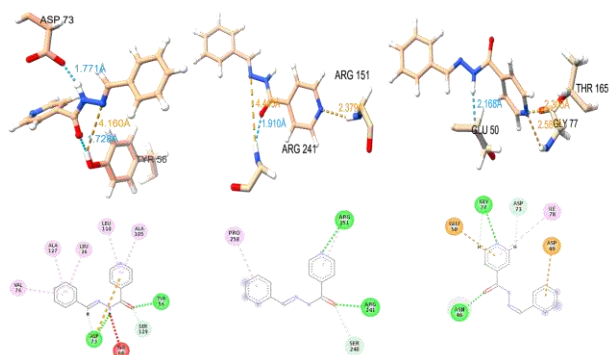


Figure 7. Docking pose of ancillary ligand against *P. aeruginosa* PDB: 3IX3, MRSA PDB: 5M18, and *E. coli* DNA Gyrase PDB: 6F86. The simulated interaction with MRSA (-6.119) displayed strong hydrogen bonding with ARG151 (2.379 Å) and ARG241 (1.910 Å). Similar interactions with *E. coli* DNA Gyrase (-6.646) and *P. aeruginosa* (-7.735) displayed essential hydrogen bonds with acidic residues such as ASP 73 (1.771 Å) and GLU 50 (2.168 Å), as well as stabilizing interactions with TYR 56 (1.728 Å).

These docking profiles provide theoretical evidence that, in addition to serving as a tuning moiety for the Mo core, the hydrazone ligand, upon its likely selective photo-release, may act as a broad-spectrum pharmacophore capable of disrupting essential pathogenic pathways. This synergistic activity makes the tricarbonyl complex a potentially viable photochemotherapeutic agent.

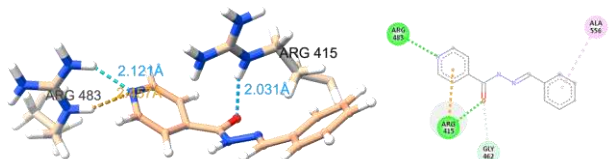


Figure 8. Binding interaction of the ancillary ligand with PDB: 4L7B.

3.7. In Vitro Antibacterial Activity

To evaluate the antibacterial potential of the complex, it was subject to in vitro susceptibility testing against *S. aureus* (Gram-positive), *E. coli* (Gram-negative), and *P. aeruginosa* (Gram-negative). The results showed potent dose-dependent zones of inhibition across all test organisms (Fig 9). The carbonyl exhibited exceptional broad-spectrum actions, yielding maximum zones of inhibition above 22 mm for the three strains. Even at the lowest concentration of 250 μM used in the evaluation, the carbonyl complex clearly outperformed the Amoxicillin control. The in vitro

susceptibility of these pathogens remarkably mirrors the CO release and delivery ability of the complex, as well as the ability of the released hydrazone to halt the growth of both Gram-positive and Gram-negative bacteria. This proves that the hydrazone is a highly effective antibacterial pharmacophore that can operate synergistically alongside liberated CO.

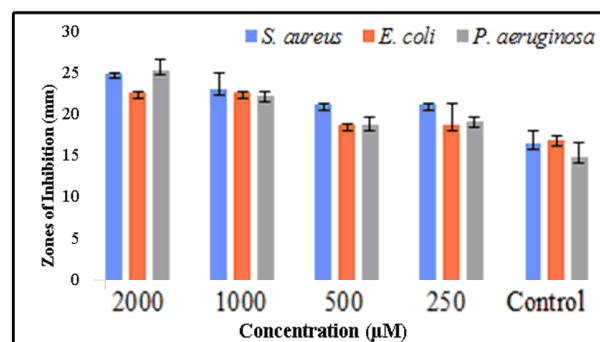


Figure 9. Antibacterial action of the CORM and Control against the test organisms.

4. Conclusion

The novel mer-[Mo(CO)₃(BIH)₃] was successfully prepared by one-pot in vacuo and characterized using spectroscopic techniques. Experimental and theoretical evaluation of its potential as a synergistic, light-triggered therapeutic was carried out. Spectral analysis revealed that the nitrogen atom in the para-position of the isonicotinoylhydrazone exerts a strong electron-withdrawing effect. This weakens π -backbonding from the Mo to the carbonyl ligands and reduces the strength of the Mo-C bond, thereby rendering the complex susceptible to rapid CO release ($k_{\text{CO}} = 9.7 \times 10^{-2} \text{ s}^{-1}$). Furthermore, TD-DFT calculations identified a robust, visible light-driven Metal-to-Ligand Charge Transfer (MLCT) transition at 587.45 nm as the primary trigger for photolysis, thereby validating the photolability of the molybdenum(0) complex.

The complex's structural design enables a dual-action therapeutic mechanism. Visible light irradiation potentially guarantees both the spatial release of cytoprotective CO and the liberation of the carefully functionalized hydrazone ancillary ligand. As validated through SwissDock modelling and in vitro assay, docking against the bacterial LasR quorum-sensing receptor of *P. aeruginosa* (PDB:31X3), PBP2a transpeptidase of MRSA (PDB: SM18), and the DNA Gyrase B subunit of *E. coli* (PDB: 6F86), as well as Keap1 (PDB: 4L78) for antioxidant response simulation, revealed that the photo-released scaffold possesses the structural prerequisite to engage critical antimicrobial and antioxidant pathways. The in

vitro testing supported the broad-spectrum action of the carbonyl. Consequently, this tricarbonyl photoCORM represents a highly promising, tunable platform for the targeted delivery of synergistic antibacterial and anti-inflammatory therapies.

Author contributions

Tenimu Adogah Abubakar: Conceptualization, methodology, software, original draft. **Abdulakeem Busari:** Writing, reviewing. **Jude Ehweverhere Emurotu:** Resources. **Dele Peter Fapojuwo:** Formal Analysis. **Oluwagbemiga T. Amusan:** Editing. **Uche Basil Eke:** Conceptualization, Reviewing, Editing, Validation, supervision.

Conflicts of interest

There are no conflicts to declare.

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