Innovators in chemistry

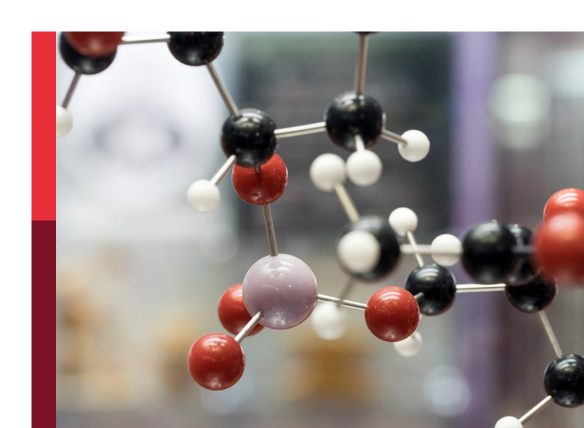
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Aleksandar Kondinski, Kunyu Wang, Jose Luis Cabellos, Arjun Singh, Barkat A. Khan, Maxence Wisztorski, Jafar Soleymani, Citlalli Gaona-Tiburcio, Tao Wei and Linghui Qian

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Editorial: Innovators in chemistry: 2022

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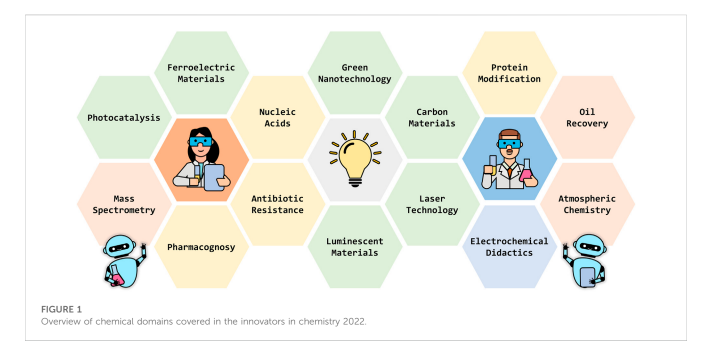
Editorial on the Research Topic

Innovators in chemistry: 2022

The "Innovators in Chemistry 2022" edition showcases the dynamic evolution and critical importance of chemistry in addressing complex challenges across various fields, emphasizing its role in driving sustainable advancements. This issue, curated by a number of young guest editors, assembles a diverse group of both established and emerging researchers whose work significantly impacts the field. Featuring 14 meticulously crafted articles by 80 researchers, the Research Topic spans a wide array of topics in chemical science, including nucleic acid structures, antibiotic resistance, pharmacognosy, green nanotechnology, photocatalysis, ferroelectric materials, protein modification, carbon materials, mass spectrometry, laser technology, luminescent materials, atmospheric chemistry, oil recovery technology, and didactical electrochemistry (Figure 1). It offers a comprehensive view of current trends and future innovations through original research, reviews and opinion articles. This edition not only celebrates recent advancements but also seeks to inspire ongoing exploration and progress in chemistry. Conclusively, "Innovators in Chemistry 2022" highlights the collaborative and intellectual contributions of its participants in steering the future of this central science. The guest editorial team expresses deep gratitude to each author, reviewer, and reader for their invaluable involvement in advancing innovative solutions and understanding of chemistry.

In this edition, we explore a number of exceptional research directions in life sciences. Monsen presents a comprehensive review on G-quadruplexes (G4s), that is four-stranded nucleic acid structures forming in guanine-rich regions of chromatin, particularly in gene promoters. His work discusses the significance of G4s as epigenetic elements in gene transcription and their potential as indirect drug targets in cancer therapies, advocating for research focused on higher-order G4s for more selective molecular targeting, using the human telomerase reverse transcriptase (hTERT) core promoter G-quadruplex as a case study (Monsen). Complementing this, Krco et al. delve into the challenge of antibiotic resistance, with a special focus on metallo- β -lactamases (MBLs) that inactivate β -lactam antibiotics. Their review emphasizes the structurefunction relationships of B3-type MBLs, noting their active site diversity and the potential for broader-range inhibitors, as suggested by the inhibition of one B3-type MBL by clavulanic acid (Krco et al.). In a distinct but equally innovative domain, Myklebust et al. report the development of potent bisubstrate inhibitors targeting NAA80, an enzyme crucial for actin N-terminal acetylation in eukaryotic cells. Their optimized compound, CoA-Ac-EDDI-NH2, combines coenzyme A and a tetrapeptide

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amide, showing high efficacy and promising insights into the roles of actin and the regulatory functions of NAA80 (Myklebust et al.). Lastly, Shaheen et al. provide a comprehensive analysis of Xanthium strumarium L. (XSL) foliage, exploring its antioxidant and anti-diabetic properties. They demonstrate that the ethyl acetate fraction of XSL, with its high phenolic concentration, significantly reduces blood glucose levels in diabetic mice, indicating a link to its traditional use in diabetes treatments (Shaheen et al.).

The landscape of advanced materials technology is rapidly evolving, as demonstrated by a series of recent studies. Zidane et al. exemplify this evolution through their work on ecofriendly nanoparticle synthesis using Purslane leaf extract, marking a significant departure from traditional hazardous chemical methods towards environmentally sustainable practices. This theme of sustainability is further explored by Lu et al., who unveil the potential of transition metal single-atom catalysts supported by red phosphorus to enhance photocatalytic hydrogen production, a crucial aspect in the search for clean energy solutions. The innovative use of materials continues in the realm of biological imaging, as Thornton et al. develop unique macrocyclic lanthanide complexes, opening new avenues in advanced imaging techniques. Expanding the boundaries of material science, Mills et al. delve into the complex effects of doping on the ferroelectric and dielectric properties of barium titanate, emphasizing the complex relationship between material composition and its properties. Concurrently, Ni et al. pose their viewpoint on the advancements in carbon-based materials, highlighting the diverse applications of porous carbon in fields ranging from adsorption and catalysis to energy storage, crucial for tackling current energy and environmental challenges. Finally, the innovative synthesis of two-dimensional transition metal dichalcogenides using laser-assisted techniques, as detailed by Wang et al., showcases a method that promises

to trailblaze flexible electronics with its precision and dynamic properties. Collectively, these studies not only reflect the current state of advanced materials technology but also chart a path for future research and applications in this rapidly evolving field.

Recent advancements in chemical analysis and control technologies are highlighted in a series of studies, each focusing on different applications and implications. Reynaud et al. present a significant leap in the field of mass spectrometry (MS) with their development of a compact nano-electro-mechanical system (NEMS) based MS, designed for ultra-high mass analytes. This prototype, operating under higher pressures without compromising on particle focusing or mass measurement quality, demonstrates an improved capability in nanoparticle analysis, a crucial advancement for MS technology. Complementing this, Dash et al. delve into atmospheric chemistry, investigating the formation of methanimine in the troposphere. By exploring the reaction dynamics of the aminomethyl radical with oxygen and considering the influences of ammonia and water, their study sheds light on the complex chemical processes occurring in our atmosphere, contributing to a deeper understanding of tropospheric compound formation. Du et al. address a practical challenge in the oil recovery industry, focusing on the flow resistance characteristics of steam injection in horizontal oil wells. Their experimental approach to evaluating pressure loss and optimizing steam distribution offers a refined method for improving oil recovery, showcasing the application of fluid dynamics and pressure control in an industrial setting.

In response to the increasing importance of electroanalytical chemistry increasing significance, Espinoza-Montero et al. developed a didactical guide on generating hydroxyl radicals ·OH using boron-doped diamond (BDD) in electrochemical oxidation processes. This educational resource elucidates key aspects such as BDD activation, electrolyte response analysis, and overpotential determination for ·OH production through Tafel plots and

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voltammetry, along with strategies for radical entrapment. Additionally, their study extends to the practical application of these methods for the efficient and cost-effective degradation of amoxicillin, thereby showcasing the distinct electronic characteristics and stability of BDD in redox reactions (Espinoza-Montero et al.).

On behalf of the involved guest editors and myself, I express our gratitude to the contributors and reviewers who played a vital role in shaping this Research Topic. Gratitude is also extended to the Frontiers team for their invaluable support, making this endeavor both enjoyable and timely. We hope this edition delights the readers of Frontiers in Chemistry and showcases the significant advancements. It is our aspiration that this Research Topic will spark interest, initiate new thoughts, and shed light on uncharted areas of chemical innovation.

Author contributions

AK: Conceptualization, Visualization, Writing-original draft, Writing-review and editing.

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Conflict of interest

Author AK was employed by Cambridge Centre for Advanced Research and Education in Singapore Ltd.

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Green synthesis of multifunctional MgO@AgO/Ag₂O nanocomposite for photocatalytic degradation of methylene blue and toluidine blue

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Introduction: In this paper, MgO@AgO/Ag $_2$ O nanoparticles were greenly synthesized, the current idea is to replace the harmful chemical technique with an ecofriendly synthesis of metal oxide nanoparticles (NPs) utilizing biogenic sources.

Methods: The current investigation was conducted to create silver oxide NPs decorated by MgO NPs (namely, MgO@AgO/Ag₂O nanocom-posite) using the leaves extract of Purslane (*Portulaca Oleracea*) as the reducing and capping agent. The nanopowder was investigated by means of X-ray diffraction, scanning electron mi-croscope, BET surface area, Fourier transform infrared, and UV-vis spectrophotom-eter studies. XRD studies reveal the monophasic nature of these highly crystalline silver nano-particles. SEM studies the shape and morphology of the synthesis AgO/Ag₂O and MgO@AgO/Ag₂O NPs. The presence of magnesium and oxygen was further confirmed by EDS profile.

Results and discussion: The surface area was found to be 9.1787 m²/g and 7.7166 m²/g, respectively. FTIR analysis showed the presence of specific functional groups. UV-vis spectrophotometer studies show the absorption band at 450 nm due to surface plasmon resonance. The results have also indicated the high performance of the greenly synthesized AgO/Ag₂O NPs and

Abbreviations: NPs, Nanoparticles; MB, Methylene blue; TB, toluidine blue.

 $MgO@AgO/Ag_2O$ NPs for photocatalytic activity dye degradation (methylene blue and toluidine blue).

KEYWORDS

MgO@AgO/Ag2O NPs, green synthesis, *Portulaca oleracea*, photocatalysis, methylene blue, toluidine blue

1 Introduction

With widespread applications in the domains of biology, health, energy, and material science, among others, nanotechnology has emerged as a promising interdisciplinary field of the twenty-first century. Nanomaterials are produced often by a variety of physical and chemical processes that call for high temperatures, vacuum conditions, specialized equipment, and chemical additives (Cai et al., 2022). Due to the use of harmful compounds that remain attached to the synthesized nanoparticles, current improvements in chemical processes for the production of nanomaterials have raised biological dangers to the environment (Yu et al., 2022). Consequently, scientists are presently concentrating on the production of nanomaterials utilizing biogenic sources such as bacteria, algae, and plants. A new age for secure nanobiotechnology has begun because of recent advancements in the green synthesis of nanomaterials, which are rapid, inexpensive, and ecofriendly. Due to their unique physical and chemical features in the domains of biosensors, diagnostic tools, catalysts, anticancer, and antimicrobial agents, metal oxide nanoparticles have recently attracted a lot of attention (Zhang et al., 2020; Ahmad et al., 2021; Wijesinghe et al., 2021a; Wijesinghe et al., 2021b; Gherbi et al., 2022).

In part because of their non-biodegradable and persistent character, the considerable water pollution caused by inorganic (such as heavy metal ions) and organic compounds—such as synthetic organic colors, persistent organic pollutants, antibiotics, etc.—often present significant obstacles. Additionally, these emerging pollutants may have several detrimental effects on human bodies and aquatic microbes (Sharma and Bhattacharya, 2017; Al-Essa and Al-Essa, 2021). The world's use of synthetic dyes now exceeds several million tons, with numerous industrial sectors including the printing and textile industries contributing to this (Mashkoor and Nasar, 2020). Therefore, solutions for wastewater pretreatment are required to reduce these contamination issues. Scientifically, the adsorption process is the most effective and practical method for purifying water is adsorption (Al-Essa, 2018; Bulgariu et al., 2019). Adsorbents should be created from free, sustainable, and locally accessible sources to achieve competitive efficacy (typically, plant resources) (Bulgariu et al., 2019).

Due to their high conductivity, silver oxide nanoparticles (AgONPs) stand out among the competition as a desirable material for metallic-based conductive filler (Stewart et al., 2017), capacity to encourage cell proliferation and osteogenic differentiation facilitate bone repair (Zhang et al., 2015; Xu et al., 2020), in addition to their capacity to prevent the development of microbial biofilm (Hileuskaya et al., 2020) and maybe defeat microorganisms that

are resistant to many drugs (Rai et al., 2012). AgONPs were highly sought-after for application in biosensors due to these special characteristics (Zeng et al., 2014; Al Essa et al., 2021), prostheses (Gallo et al., 2016), and dental materials (Ohashi et al., 2004). The resultant AgONPs, however, are inappropriate for biomedicine use due to the common synthesis technique, which frequently employs toxic chemical reducing agents that are dangerous for human (such as sodium borohydride and hydrazine) (Kolarova et al., 2015; Rónavári et al., 2021). Magnesium oxide nanoparticles (MgONPs) are harmless and reasonably simple to synthesize among the many distinct inorganic metal oxides. MgONPs have received approval from the United States Food and Drug Administration as safe materials (21CFR184.1431) (Martinez-Boubeta et al., 2010). MgONPs, for instance, can reduce indigestion, start the postactivation of bone-repair scaffolds, and function as hyperthermia agents in cancer treatment. Both MgO and AgO NPs can be now produced using "green synthesis" techniques, which offer straightforward, affordable, quick, and non-toxic approaches to address these problems. The use of diverse biological entities, such as plant extract, in green synthesis techniques replaces the use of chemical reducing agents. (Ahmadi et al., 2018; Hamouda et al., 2019; Okaiyeto et al., 2019), and bacteria (Hamouda et al., 2019) produce a high output of somewhat uniform-sized metallic nanoparticles by acting as both reducing and capping agents. Due to their accessibility, safety, and simplicity of synthesis, plant-based extracts have drawn more attention. They don't require the preservation of microbial cultures (Roy and Das, 2015). The most often reported use so far has been of various plant fruit and leaf extracts. Purslane is a fascinating possibility for this use among many different plants (Bouafia et al., 2020; Bouafia et al., 2021; Bouafia and Laouini, 2021).

The decor is another method for enhancing photocatalytic activity (Wang et al., 2011; Hunge et al., 2021; Yadav et al., 2021). Surface plasmon resonance (SPR), where Ag, Au, and Pt are typically utilized, is favored by decoration with noble metals (Xing et al., 2017), it has many positive benefits since it promotes the absorption of visible light and the activation of energetic charge carriers, many (Sohrabnezhad and Seifi, 2016), achieves a greater separation of charge, this leads to an increase in redox reactions and a higher formation rate of reactive oxygen species (Ong et al., 2013; Kuriakose et al., 2014; Sohrabnezhad and Seifi, 2016). Previous studies have confirmed the photocatalytic activity of AgO and Mg NPs (Balakrishnan et al., 2020). Moreover, prior studies demonstrated that the decoration of AgONPs with other metals NPs results in a significant improvement in the photocatalytic activity comparing to the nondecorated AgONPs (Laouini et al., 2021).

In this study, the degradation of the methylene blue (MB) and toluidine blue (TB) dyes under sunlight was used to examine the photocatalytic capabilities of ${\rm AgO/Ag_2O}$ nanoparticles (NPs) coated with MgO. To the best of our knowledge ${\rm AgO/Ag_2O}$ and MgO@AgO/Ag₂O NPs photocatalytic activity was never tested against MB and TB dyes.

2 Materials and methods

2.1 Reagents

Silver nitrate (AgNO₃, 98%), Magnesium chloride hydrates (MgCl₂, $6H_2O$, 98%), MB ($C_{16}H_{18}ClN_3S$, 98%), and TB ($C_{15}H_{16}ClN_3S$, 98%) were purchased from Sigma-Aldrich,Germany. The leaves of purslane were collected from local fields in El Oued, Southeast of Algeria. Distilled water was used in all the experiments.

2.2 Preparation of the leaf extract

Fresh and healthy purslane (*Portulaca oleracea*) leaves were collected from nearby farms in the El Oued area (Southeast of Algeria). After properly cleaning them with flowing tap water to get rid of any dirts or other polluted organic materials, they were repeatedly rinsed in de-mineralized water. The fresh leaves were crushed. The leaves components were extracted by mixing 250.0 g of leaves with 900 ml of distilled water in a 1,000 ml glass beaker. The mixture was stirred for 15 min at 70°C. The extract was kept cool and then filtered using the decantation method and stored at 4°C for further use.

2.3 Synthesis of AgO/Ag₂O and MgO@ AgO/Ag₂O NPs

The AgO/Ag₂O and MgO@AgO/Ag₂O NPs were synthesized by a green method using plant extract, following modified protocols from previous studies (Abdullah et al., 2020; Bouafia and Laouini, 2020; Bouafia et al., 2021a; Bouafia et al., 2021b; Belaiche et al., 2021; Laid et al., 2021; Laouini et al., 2021; Ben Amor et al., 2022; Daoudi et al., 2022; Djamila et al., 2022; Meneceur et al., 2022; Tedjani et al., 2022). For the synthesis of AgO/Ag₂O NPs filtered extract was taken and diluted to 1,000 ml with deionized water, 1.7 g of AgNO₃ was added to 1,000 ml of the extract with continuous stirring, 5 ml diluted hydroxide (0.1 M) was added dropwise to the mixture solution. With controlled and continuous stirring (600 rpm) at 70°C for 4 h, the precursor became brown. The precipitate was then centrifuged and washed several times with deionized water, and dried at 100°C in an oven for 48 h. Finally, the AgO/ Ag₂O NPs were manually grinded into a fine powder.

Nanostructured MgO loaded AgO/Ag₂O were simply synthesized. Firstly, nanoparticle solution (AgO/Ag₂O) was prepared by dissolving 0.7 g of (AgO/Ag₂O) in de-ionized water, the solution maintained in continuous stirring at 70° C for 2 h. Then 70 ml (0.01 M) metal ion (MgCl₂) solution was added to the nanoparticle solution, mixed thoroughly at 80° C for 2 h, and then dried at 100° C for 48 h.

After drying, the mixture was grinded to obtain a homogeneous powder.

2.4 Material characterizations

The employment of the following methods was necessary for the characterization of generated nanoparticles: X-ray diffractometer (XRD), a scanning electron microscope (SEM), EDS, BET surface area analyzer, Fourier transform infrared (FTIR), and UV-vis spectrophotometer.

Using XRD (Rigaka Miniflex 600) and Cu-K radiation with a wavelength of 0.15406 nm in the 2 range 10-80, the crystalline structure of the produced NPs was determined. The morphology of the prepared AgO/Ag2O and MgO@AgO/Ag2O NPs were verified using SEM (TESCAN VEGA 3) with an accelerating voltage of 10 kV. The BET surface area analyzer (micromeritics ASAP 2020 Plus Version 2.00) was used to capture the N2 adsorption isotherm, associated surface area, and pore parameter. A Nicolet iS5 (Thermo Fisher Scientific) was used to perform FTIR measurements on leaves extract, green synthesis AgO/Ag₂O, and MgO@AgO/Ag₂O NPs to determine the functional groups operating in the region of 4,000 to 400 cm⁻¹. A UV-vis spectrophotometer was used to examine the optical properties of the samples (Shimadzu-1800). The measurement was made at the temperature of 28°C in the 300-900 nm wavelength range. UV-vis spectrometric measurements were performed using a quartz cell and distilled water as a blank solution to determine the stability of AgO/Ag₂O NPs. The optical gap band energy (Egap) of these materials was calculated using the Tauc Equation considering these findings.

2.5 Photocatalytic degradation of MB dye

Based on the samples' capacity to break down the MB dye in the presence of sunlight, the catalytic activity of the samples was evaluated. The experiment was conducted in a lab using sunlight (the samples were not exposed to direct sunlight). MB stock solution ($2.5 \times 10^{-5}\,\mathrm{M}$) was made. Four samples of 5 ml of dye solution and 5.0 mg of AgO/Ag₂O NPs were combined and exposed to sunlight in. About 2 ml of the suspensions were taken from the reaction mixture every 15, 30, 60, and 120 min while it was exposed to sunlight, and the suspended particles were then extracted using ultracentrifugation. The same test was

conducted using MgO@AgO/Ag $_2$ O. The absorption obtained at 664 nm in a UV-vis spectrophotometer was used to calculate the rate of dye degradation. Based on the formula, the dye degradation % was evaluated using the Eq. 1:

% of degradation =
$$\frac{C_i - C_t}{C_t}$$
 (1)

where C_t and C_t (mg/L) are the initial concentration of MB and the concentration of pollutant MB at time t, respectively.

2.6 Photocatalytic degradation of TB dye

The photodegradation of TB dye in aqueous solution was investigated under sunlight in the presence of AgO/Ag₂ONPs and MgO@AgO/Ag₂O as a photocatalyst, A5.0 mg of the catalyst is added to 50 ml of TB dye solution. This solution is prepared in distilled water at a concentration of (4 \times 10 $^{-3}$ M). It was then exposed to sunlight at different intervals (10, 20, 30, 40, and 60 min) at a temperature of 28°C and a neutral pH. The catalyst is separated by ultracentrifugation. A spectrophotometer is used for reading the absorbance values at 631 nm.

The equilibrium amount of adsorption is calculated by the Eq. $\stackrel{\hbox{\scriptsize 2}}{}$:

$$QE = \frac{(C_0 - C_e)V}{m}$$
 (2)

where, QE (mg/g) is the adsorption capacity at equilibrium, C_0 and C_e (mg/L) are the concentration of TB solutions at the start of reaction and the concentration of pollutant TB at time t, respectively. While V stands for the volume of the solution (L), and m is the mass of adsorbent (g).

3 Results and discussion

3.1 Crystal structure and composition

Portulaca oleracea's medicinal potency is attributed to its complex chemical makeup composition; it is high in primary and secondary metabolites, as well as minerals, vitamins, and other micronutrients.

Purslane is the common name for *Portulaca oleracea L.*, a member of the *Portulaceae* family. A variety of substances have been identified from *Portulaca oleracea*, including flavonoids, alkaloids, polysaccharides, fatty acids, terpenoids, sterols, proteins, vitamins, and minerals. There is a wide range of pharmacological qualities that *Portulaca oleracea* exhibits, including neuroprotective, antibacterial, antidiabetic, antioxidant, anti-inflammatory, antiulcerogenic, and anticancer actions. (Zhou et al., 2015)

To investigate the size and make-up of the crystalline phases, XRD analysis was done.

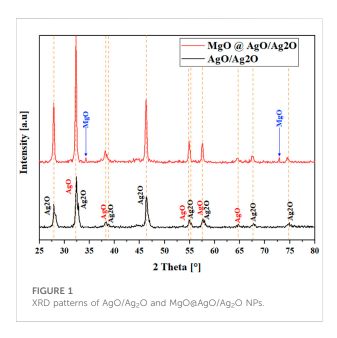


Figure 1 shows the diffraction patterns of AgO/Ag₂O and MgO@AgO/Ag₂O. Similar results have already been described in the literature (Deekshitha and Shetty, 2021), regarding the peaks of Ag₂O. Moreover, It is intriguing to see that there are two distinct peaks that can be attributed to MgO NPs. According to the Debye-Scherrer equation, the crystallite size was determined to be 22.16 nm for AgO/Ag₂O and 21.60 nm for MgO@AgO/Ag₂O NPs, respectively. The results are shown below in Table 1.

3.2 Morphological investigation

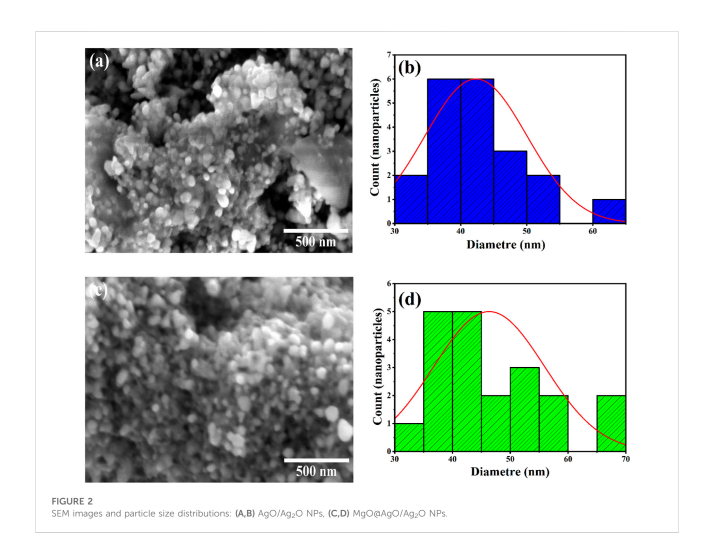
SEM was used to study the formation of the prepared AgO/Ag₂O and MgO@AgO/Ag₂O NPs (Figures 2A,C, respectively), and their morphological size (40–50 nm in average) (Figures 2B, D, respectively). The MgO@AgO/Ag₂O NPs were oval and spherical. Most of the MgO@AgO/Ag₂O NPs were placed as aggregated which may explain the slight increase in size of these NPs compared to AgO/Ag₂O NPs; also, a few individual particles were also observed (Gosens et al., 2010).

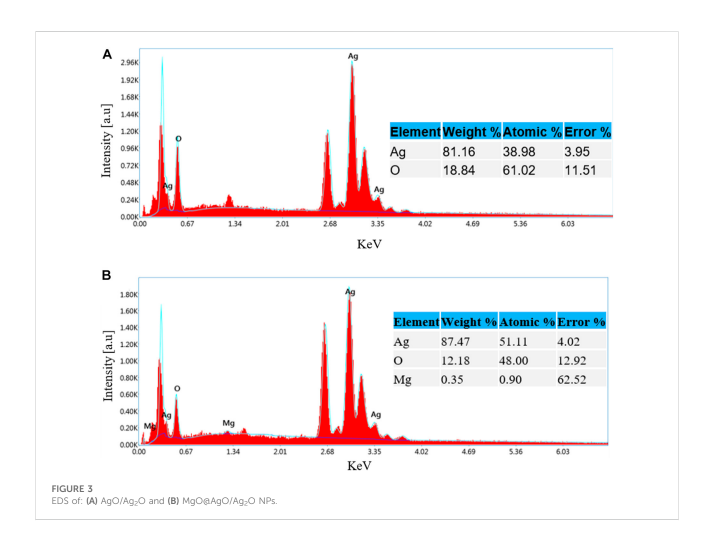
Further analysis by EDS of the prepared AgO/Ag₂O and MgO@AgO/Ag₂O NPs (Figures 3A,B, respectively). The data associated to MgO@AgO/Ag₂O NPs, confirm the presence of silver, oxygen, and magnesium, with a weight percentage of approximately 87.47% Ag, 12.18% O and 0.35% Mg. It is concluded that all nanoparticles were mixed together. In this way, the aimed ternary oxides nanoparticle system containing MgO, AgO, and Ag₂O NPs was obtained.

The MgO@AgO/Ag₂O NPs sample's EDAX analysis revealed only clear peaks for the elements Ag, Mg, and O; no additional peaks could be observed, proving that the powder was produced without any impurities.

TABLE 1 Quantification and average crystallite size of AgO/Ag $_2$ O and MgO@AgO/Ag $_2$ O NPs.

		-					
Crystallite size (nm)	Presented Phases			Lattice parameters	Crystal system	COD Entry	References
	Name	Amount (%)	Formula		3,3,0,111	Littiy	
21.60	Magnesium Oxide Periclase	31.1	MgO	Space Group: Fd-3m E a = 8.12 Å α = 90.0000° b = 8.12 Å β = 90.0000° c = 8.12 Å γ = 90.0000°	Cubic	96-500- 0226	Freund and Farmer, (1974)
	Silver oxide	15.3	AgO	Space Group: P 1 (#1-1) $a = 4.21 \ \mathring{A} \ \alpha =$ 90.0000° $b = 4.21 \ \mathring{A} \ \beta = 90.0000$ ° $c = 4.21 \ \mathring{A}$ $\gamma = 90.0000$ °	Cubic	01-076- 1489	Stehlik et al. (1959)
		55.7	Ag ₂ O	Space Group: P 1 (#1-1) $a = 4.71 \ \mathring{A} \ \alpha = 90.0000^\circ \ b = 4.71 \ \mathring{A} \ \beta = 90.0000 \ c = 4.71 \ \mathring{A}$ $\gamma = 90.0000^\circ$	Cubic	00-003- 0796	Tulsky and Long, (2001)
22.16	Silver oxide	22%	AgO	Space Group: P 1 (#1-1) $a = 4.21 \ \mathring{A} \ \alpha = 90.0000^\circ \ b = 4.21 \ \mathring{A} \ \beta = 90.0000 \ c = 4.21 \ \mathring{A}$ $\gamma = 90.0000^\circ$	Cubic	01-076- 1489	Stehlik et al. (1959)
		78%	Ag ₂ O	Space Group: P 1 (#1-1) $a=4.71 \ \mathring{A} \ \alpha=90.0000 \ b=4.71 \ \mathring{A} \ \beta=90.0000 \ c=4.71 \ \mathring{A}$ $\gamma=90.0000^\circ$	Cubic	00-003- 0796	Tulsky and Long, (2001)





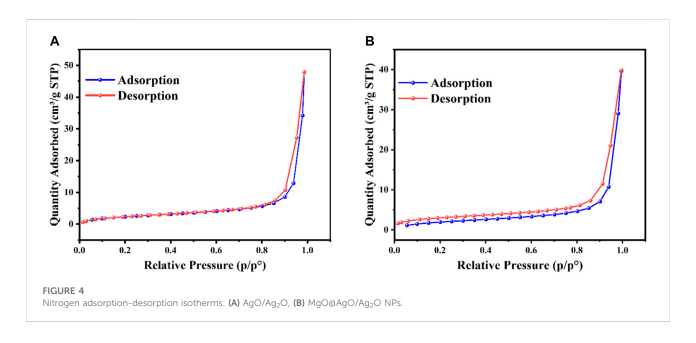
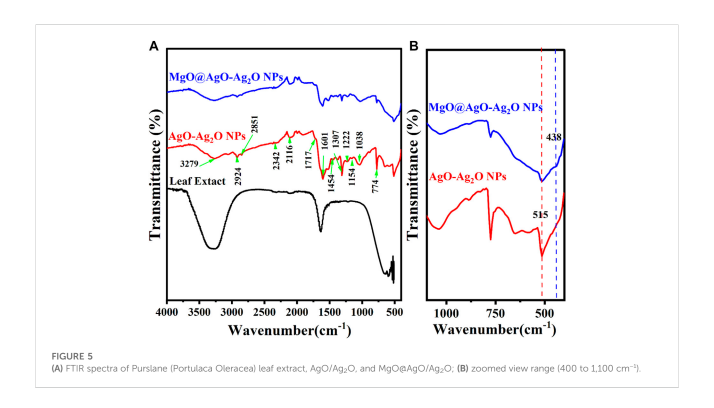


TABLE 2 BET surface area, porosity, and particle size of AgO/Ag₂O and MgO@AgO/Ag₂O NPs.

	Samples	AgO/Ag ₂ O	MgO@AgO/ Ag₂O
Surface Area	BET Surface Area	9.1787 m²/g	7.7166 m ² /g
	Langmuir Surface Area	376.2072 m²/g	279.6492 m²/g
Pore Volume	Single point adsorption total pore volume of pores less than 40.3122 nm diameter at $p/p^{\circ} = 0.950000000$	0.021557 cm ³ /g	0.020119 cm ³ /g
	Single point desorption total pore volume of pores less than 40.3122 nm diameter at $p/p^{\circ} = 0.950000000$	0.039853 cm ³ /g	0.033269 cm ³ /g
Pore Size	BJH Adsorption average pore diameter (4V/A)	3.10194 nm	3.15833 nm
	BJH Desorption average pore diameter (4V/A)	2.89018 nm	2.83265 nm
Nanoparticle Size	Average Particle Size	65.3689 nm	77.7541 nm



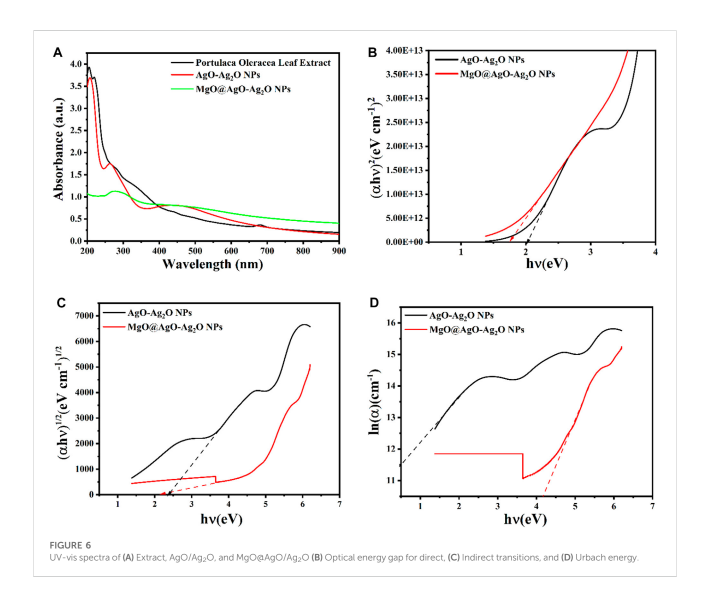
3.3 N₂ adsorption-desorption isotherm

Figure 4 shows the N_2 adsorption-desorption isotherm of 1) AgO/Ag₂O and 2) MgO@AgO/Ag₂O NPs. The specific BET surface area was found to be 9.1787 m²/g and 7.7166 m²/g (Table 2). The average pore diameter is 3.1 nm, which demonstrates the existence of mesopores. The pore size between 2.89018 and 2.83265 nm shows that the AgO/Ag₂O and MgO@AgO/Ag₂O NPs, belong to mesoporous materials (2–50 nm), this result can be easily verified by SEM images (Figure 4). The noticeable improvement, in the specific surface

area value, proves the enormous photocatalytic activity of the MgO@AgO/Ag $_2$ O NPs.

The figure below presented (Figure 4) depicts the nitrogen adsorption isotherms on the surface of the AgO/Ag_2O and $MgO@AgO/Ag_2O$ NPs. The figures clearly show that:

- All compounds of these nanoparticles belong to mesoporous materials.
- The compound MgO@AgO/Ag₂O NPs presents the best catalytic behavior with concerning the adsorption of N₂.



3.4 FTIR study

As a result of the aliphatic C-H stretching vibration of hydrocarbon chains and N-H bending vibration, the spectral peaks at 2,924, 2,851, and 1,454 cm¹⁻ (Ananthi et al., 2018; Hamouda et al., 2019) are shown in Figure 5.

Peaks centered at 1,340, 1,224, 1,142, and 1,024, and cm⁻¹ in the range 1,300–1,000 cm⁻¹ are caused by stretching vibrations of C-O groups in anhydrides, esters, ethers, alcohols, and phenols, C-O-H groups in alcohols and phenols, and C-N groups in amines. The vibration of the C=C group of alkenes, the N-H groups of amides, & the amine salts is represented because of the stretch that was seen at 1,603 cm⁻¹. It is possible that the broad stretch in the 3,400–2,400 cm⁻¹ range, which is centered at 3,283 cm⁻¹, is caused by the stretching vibrations of amines, amine salts, sulfonamides, alkenes and alkanes, amines' N-H and C-H, and carboxylic acids' C=O. Nitriles, aromatic rings, and

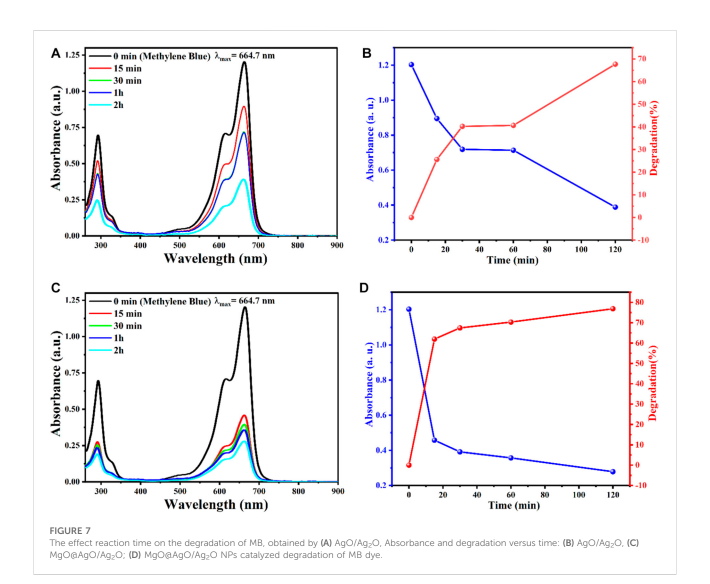
aldehydes may all be appeared as other weak bands on the graph Thus, the results of the FTIR investigation show that the main contributing factors to the reduction of Ag⁺ ions to Ag⁰ nanoparticles in the floral extract were the -C=O (carboxyl), -OH (hydroxyl), and N-H (amine) groups (Chekuri et al., 2015). The interaction of these functional groups with the AgONPs may be responsible for the shifting of these peaks (Hamouda et al., 2019).

The FTIR spectra show that the Mg-O stretching frequency of MgO corresponds to the IR peak at 436 cm⁻¹ (Sahoo et al., 2020).

Additionally, the disappearance of the phenolic compound-associated absorbance bands 3,278, 2,924, 2,851, and 1,454 cm $^{-1}$ after the synthesis of MgO@AgO/Ag2O NPs leads us to conclude that the *Portulaca oleracea L.* leaf extract contains phytochemicals like alcohols, aldehydes, alkanes, epoxy groups, and ether groups that may be responsible for the nucleation process to reduce precursor from $M^{\scriptscriptstyle +}$ to $M^{\scriptscriptstyle 0}$.

TABLE 3 Direct, indirect optical band gaps, and Urbach energies of synthesized NPs.

Samples	Urbach energy (eV)	Direct optical bandgap (eV)	Indirect optical bandgap (eV)
AgO/Ag ₂ O	0.379	2.02	2.35
MgO@AgO/Ag ₂ O	0.666	1.75	2.17

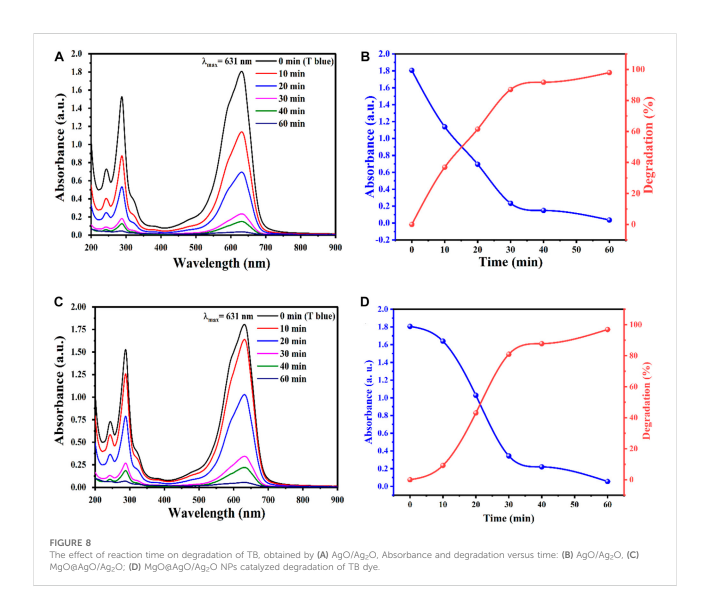


3.5 Optical properties

The formation of AgONPs was achieved within 90 min after adding 1.7~g of AgNO $_3$ to the stirred and heated extract. The process did not require the addition of external additives. In addition, bioactive molecules in leaf extracts interact with silver ions (Ag+) and fuse neighboring small particles into nanoparticles, leading to nucleation of (Ag $^{\circ}$) atoms and promoting the nanoparticle growth (Sangar et al., 2019). The formation of AgONPs was confirmed visually by the color

change of the reaction mixture from pale green to dark brown. UV-vis spectral scans are commonly implemented to confirm the formation of metal nanoparticles in aqueprobleous solution. Similarly, the product obtained exhibited a surface plasmon resonance (SPR) peak at 240 nm, as a characteristic of AgONPs (Aisida et al., 2019) (Figure 6A). Purslane extract exhibited a peak at approximately 240 nm attributed to the properties of bioactive reducing agent molecules.

To further improve the properties, we synthesized hybrid nanostructures MgO@AgO/Ag2O containing 10 mM



MgCl₂. Magnesium incorporation into the AgO/Ag₂O NPs solution was confirmed by UV-vis spectral analysis. However, the reduced SPR peak intensity is related to confirmation of MgO capping on the AgNP surface

The optical band gaps of the biosynthesized NPs were estimated using the absorption spectra of both direct and indirect transitions. The Tauc's equation (Eq. 3) was used to obtain the optical bandgap (Eg) of AgO/Ag_2O and $MgO@AgO/Ag_2O$ NPs (Soltan et al., 2017; Lassoued et al., 2018).

(Sathiyaseelan et al., 2020).

$$(\alpha h \nu) = A \Big(h \nu - E_g^{opt} \Big)^n \tag{3}$$

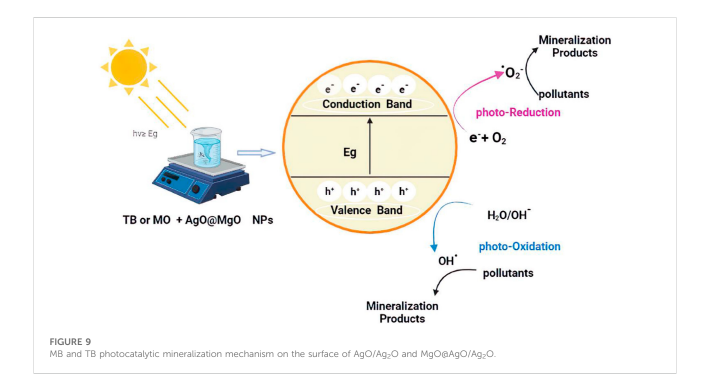
According to Lambert-Beer-Bouguer Law, the absorption coefficient (Eq. 4) is given as (Kilic et al., 2019):

$$\alpha = \frac{A}{t} \ 2.303 \tag{4}$$

here, a represents the absorbance and t is the of quartz cuvette width (10 mm).

For the direct transition Tauc plots were plotted using $(\alpha h v)^2$ against hv, meanwhile $(\alpha h v)^{V2}$ against hv were used to plot the indirect transition graph for each sample. Were hv, h, v and α represents the photon energy, Planck's constant, the photon frequency, and the absorption coefficient respectively.

The indirect optical bandgap values of the biosynthesized AgO/Ag₂O and MgO@AgO/Ag₂O NPs varied from 2.35 to 2.17 eV (Figure 6C), while the direct bandgap values varied from 2.02 to 1.75 eV (Figure 6B). The direct optical bandgap of the synthesized sample increased with decoration, while the indirect optical bandgap decreased. Higher bandgap values may indicate the presence of confinement effects in the manufactured product. Such high bandgap values of prepared nanostructures make them suitable for electro-optical devices (Shakir et al., 2009; Shkir et al., 2018).



The Urbach energies (Eq. 5) can be estimated exploiting the slope of the absorption edge in the semi-logarithmic plot (Figure 6D).

$$\ln \alpha = \frac{hv}{E_n} + \text{constant } (\ln \alpha_0)$$
 (5)

here, E_u signifies the Urbach energy.

During our investigation, we observed changes in the Urbach energy of the biosynthesized NPs. These changes occurred due to the influence of MgO. Where an increase in the Urbach energy (from 0.379 to 0.666 eV) was observed upon MgO incorporation, indicating a distortion of the structural order. The Urbach energy characterizes the homogeneity and stability of NPs. The lower the Urbach energy, the more uniform and stable the nanoparticles, and *vice versa*. The increasing AgO addition to the glass structure explains that the structure has become disordered and unstable. This result is consistent with the literature (Table 3) (Işsever et al., 2019).

3.6 Photocatalytic activity of AgO/Ag₂O and MgO@AgO/Ag₂O for azo dye degradation

The organic MB dye was used to measure the catalytic activity of AgO/Ag_2O and $MgO@AgO/Ag_2O$ NPS by measuring the decomposition rate of MB. Selected at 664 nm.

The percentage of MB degradation by AgO/Ag_2O and $MgO@AgO/Ag_2O$ NPs in the presence of sunlight is shown in

(Figures 7B, D). The results showed that the percentage MB decomposition values of synthesized AgO/Ag $_2$ O and MgO@ AgO/Ag $_2$ O NPs changed from 67.71363% to 76.86329% in the presence of sunlight.

According to (Figures 7A, C), magnesium decoration had a significant effect on the photocatalytic efficiency of AgO/Ag₂O powders. The photocatalytic efficiency of AgO/Ag₂O powder was increased by magnesium decoration. As it can been seen, the photocatalytic efficiency was directly proportional to the exposure duration (time).

This result is caused by surface plasmon resonance (SPR), resulting in increased redox reactions and a higher rate of reactive oxygen species generation. Additionally, heterojunctions structured between AgO/Ag₂O and MgO@AgO/Ag₂O NPs act to increase charge carrier separation, thereby providing highly reactive species that function in photocatalysis (Wang et al., 2011; Ong et al., 2013; Paula et al., 2019).

The effect of photocatalyst decomposition is shown in the Figure 8. Irradiation of NPs affects the degradation of pollutant TB. This effect may be due to the availability of sufficient energy to excite electrons from the valence band to the conduction band. The band gap energy values for AgO/Ag₂O and MgO@AgO/Ag₂O NPs (2.02 eV to 1.75 eV) justify the reduction of the band gap energy. The reaction mechanism is shown in Figure 9.

Wastewater purification remains a major environmental issue due to the increasing and widespread spread of non-degradable pollutants in water, Therefore, this study presents the optimization of photocatalytic degradation of TB dye in an aqueous medium by photocatalyst, under sun irradiation. The

obtained results revealed that most of the TB were removed within 60 min at a rate of 98%.

Photolysis is a chemical reaction in which a chemical substance is broken down by photons, whereas photocatalysis is defined as the acceleration of a photoreaction in the presence of a catalyst. Surface chemistry peculiar to this family of chemicals influences how electron donors and acceptors interact with metal oxide semiconductors. Light is absorbed by an absorption substrate during catalysed photolysis. For electrons to be promoted from the valence band (VB) to the conduction band (CB), forming electron-hole (e⁻/h⁺) pairs, photocatalyst on semiconducting oxides must absorb photons with energy equal to or greater than the oxide's band gap (Yao et al., 2008; Mittal et al., 2009), as shown in the Eq. 6.

The following steps sum up the mechanism of oxidation for the photodegradation process of the pollutant MB and TB:

Step 1: The radical anion is formed Eq. 6:

$$e_{(CB)}^{-} + O_2 \xrightarrow{AgO/Ag2O \text{ or } MgO@AgO/Ag2O} O_2^{-}$$
 (6)

Step 2: The OH radical group appears by the hole (7-11);

$$h_{(VB)}^+ + H_2O \xrightarrow{AgO/Ag2O \text{ or } MgO@AgO/Ag2O} H^+ + OH^-$$
 (7)

$$H^+ + \dot{O}_2^- \longrightarrow H\dot{O}_2$$
 (8)

$$H\dot{O}_2 + H\dot{O}_2 \longrightarrow H_2O_2 + O_2$$
 (9)

$$H_2O_2 \xrightarrow{hv} 2O\dot{H}$$
 (10)

Step 3: The organic contaminants are oxidized either through direct contact with the holes or by the main radicals and OH (Eq. 11);

Toluidine blue or Methylene blue
$$+ OH \xrightarrow{h^+}$$
Mineralisation products (11)

4 Conclusion

The catalytic activity of AgO/Ag₂O and MgO@AgO/Ag₂O NPs was evaluated based on the rate of degradation of organic dye pollutant. Experimental data illustrated that the percentage of MB degradation values of the synthesized AgO/Ag₂O and MgO@AgO/Ag₂O NPs changed from 67.71 to 76.86%, and 98% of TB degradation. Furthermore, under environmental circumstances, the synthesized AgO/Ag₂O and MgO@AgO/Ag₂O NPs display strong photocatalytic activity for dye

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degradation of MB and TB stains. AgO/Ag₂O and MgO@ AgO/Ag₂O NPs have been found to be useful in the treatment of wastewater (dye degradation). It has proven its ability to purify water from all suspended impurities and remove chemical dyes, and there is a possibility to apply photocatalytic methods to purify water contaminated with hydrocarbons.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

Author contributions

Conceptualization, writing—original draft preparation: YZ, SL, AB, SM, MT, and FM; methodology, validation, data curation: YZ, SL, AB, SM, MT, SA, HA, KE, EMA-E, MR, OM, and FM; investigation: AB, MT, SM and FM; resources: YZ, AB, AA, KE, EMA-E, MR, and FM; writing—review and editing: AB, SM, MT, SA, HA, KE, EMA-E, MR, and FM; supervision: SL, SM, and FM; Submission: FM. The authors have read and agreed to the published version of the manuscript.

Conflict of interest

Author FM was employed by the company Fluorotronics, Inc.-California Innovations Corporation.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Laser-assisted synthesis of two-dimensional transition metal dichalcogenides: a mini review

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The atomically thin two-dimensional (2D) transition metal dichalcogenides (TMDCs) have attracted the researcher's interest in the field of flexible electronics due to their high mobility, tunable bandgaps, and mechanical flexibility. As an emerging technique, laser-assisted direct writing has been used for the synthesis of TMDCs due to its extremely high preparation accuracy, rich light-matter interaction mechanism, dynamic properties, fast preparation speed, and minimal thermal effects. Currently, this technology has been focused on the synthesis of 2D graphene, while there are few literatures that summarize the progress in direct laser writing technology in the synthesis of 2D TMDCs. Therefore, in this mini-review, the synthetic strategies of applying laser to the fabrication of 2D TMDCs have been briefly summarized and discussed, which are divided into top-down and bottom-up methods. The detailed fabrication steps, main characteristics, and mechanism of both methods are discussed. Finally, prospects and further opportunities in the booming field of laser-assisted synthesis of 2D TMDCs are addressed.

KEYWORDS

laser, two-dimensional materials, transition metal dichalcogenides, synthesis methods, mechanism

1 Introduction

Flexible electronics is a disruptive science and technology based on a high degree of interdisciplinary integration, which provides novel opportunities for the development of the next generation of the information technology revolution and the era of intelligent manufacturing (Zheng et al., 2022; Li et al., 2023). The two-dimensional materials, especially transition metal dichalcogenides have become trending research topics of flexible electronics due to their unique structure, mechanical flexibility, tunable bandgaps, and high mobility (Mak et al., 2010; Fan et al., 2015; Kim et al., 2015; Leong, 2020; Yu et al., 2020; Liu et al., 2021; Wu et al., 2022).

Currently, significant research has been focused on the efficient synthesis of high-quality large-scale 2D TMDCs with simple crafts and low costs, owing to the prerequisite and key factor for their diverse advanced industrial application (Lee et al., 2012; Lei et al., 2013; Mahjouri Samani et al., 2014; Shim et al., 2014; Xia et al., 2014; Xu et al., 2021b). Various synthesis techniques, such as chemical vapor deposition (CVD), metal-organic chemical

TABLE 1 Summary of preparation methods of 2D TMDCs.

Method	Advantage	Disadvantage
Mechanical exfoliation	High quality, simple	Low yield, challenging to mass production
Liquid-phase exfoliation	High preparation efficiency, simple	Small lateral size, low yield
Physical vapor deposition	High quality, low cost	Low repetition rate, complex process
Atomic layer deposition	High repetition rate	High cost, complex process
Hydrothermal	High yield, low cost	Low controllability, unstable grain size, low quality
CVD	Low cost, high quality, possible to mass production	High preparation temperature, low repetition rate
MOCVD	High deposition rate, high quality	Chemical pollution, high cost, long production cycle
MBE	Atomic-level control, high purity	Rigorous conditions, expensive equipment
Laser-assisted synthesis	Low cost, high repetition rate, patternable	Hardly achieve nanometer-level resolution, numerous defects

vapor deposition (MOCVD), and molecular beam epitaxy (MBE), have been used to control the growth of 2D TMDCs on different substrates and show great potential in growing high-quality crystalline 2D TMDCs under high temperature (Yu et al., 2017; Gao et al., 2021). However, patterning TMDCs cannot be achieved in the synthesis process, and an additional patterning process is still required during the device fabrication (Mak et al., 2010). In recent years, direct laser writing technology has been widely used in the synthesis of materials because of the ability to minimize thermal effects, rich light-matter interaction mechanism, kinetic properties, the high fabrication accuracy, and fast preparation speed (Lu et al., 2014; Jung et al., 2019; Park et al., 2020a; Park et al., 2020b; Hu et al., 2020). In addition, maskless and lithography-free properties of laser patterning can enable one-step fabrication of the desired specific patterns, avoiding material contamination while reducing the process flow (Cao et al., 2013; Mohapatra et al., 2020). All preparation methods of synthesizing 2D TMDCs and their advantages and disadvantages are shown in Table 1, which shows the salient features of direct laser writing technology from multiple perspectives.

In this mini-review, we systematically summarize the latest studies on the laser-assisted synthesis of 2D TMDCs. Basically, there are essentially two classes of synthetic strategies: top-down and bottom-up methods. With regard to top-down methods, the advantages and mechanisms of the laser-material interactions are discussed, including laser exfoliation, laser thinning, and laser-driven phase transition. With regard to bottom-up methods, the growth mechanisms are discussed, including direct laser writing, laser heterostructures, and laser doping. Finally, prospects and further opportunities in the laser-assisted synthesis of 2D TMDCs are also addressed.

2 Top-down synthesis methods

Top-down synthesis methods generally require synthesizing the 2D TMDC crystal first, followed by further processing via laser–material interactions to improve the morphology or properties. Specifically, it can be classified into laser stripping, laser thinning, and laser-driven phase transition.

2.1 Laser exfoliation

The unique characteristics of the laser, such as ultra-high-peak power density and short pulse duration, far exceed conventional heating methods, such as electric or gas heating, thus making it possible to exfoliate monolayers of two-dimensional materials. The van der Waals interaction between the layers of TMDC materials can be broken by the laser, which enables the efficient and controllable preparation of TMDCs with a specific number of layers in a relatively short time. In addition, this technique has now been used to exfoliate a variety of 2D TMDC materials such as MoS₂, MoSe₂, and WS₂ under different conditions (Schuffenhauer et al., 2005; Hu et al., 2006; An et al., 2018; Gao et al., 2019; Zhai et al., 2021; Zuo et al., 2021; Kimiagar and Abrinaei, 2023).

TMDCs are layered materials that form an MX2 crystal structure composed of strongly covalently bonded X-M-X sandwiches, and the X-M-X layers are held together by weak van der Waals forces (An et al., 2018). Zhai et al. (2021) created a method for exfoliating 2H-phase MoS₂ (2H-MoS₂), that is, both efficient and free from contaminants, and also the exfoliation process can be directly observed by optical microscopy (Figure 1A). Compared to other exfoliation techniques, this method is significantly faster and easier. The thickness of the irradiated region underwent a reduction approximately from 100 nm to 5 nm, and the area of the peeled region is also much larger than the diameter of the laser spot used. They believed that the light illumination and water medium are essential for the laser exfoliation of 2D TMDCs. When 2H-MoS₂ layers are exposed to laser irradiation, the rapid vaporization of water molecules between the layers results in the exfoliation of MoS₂ sheets, and the exfoliation process can be controlled by applying a bias voltage. The related research results indicated that water molecules could be incorporated into the interlayer spaces of 2H-MoS2 sheets (Levita and Righi, 2017; Ma et al., 2018). The phase transition may accompany the exfoliation process. Gao et al. (2019) achieved a one-step exfoliating bulk 2H-MoS2 into 2H and 1T MoS2 nanosheets using pulsed laser irradiation. Figure 1B shows a conceptual model of this experimental mechanism, suggesting that the 2H-phase MoS₂ can be reversed into 1T MoS₂ with the assistance of the chemically doped Fe³⁺ ions. In addition, the capability of the protic solvent also provides the proton to the reaction environment and plays a crucial role in triggering the phase transition.

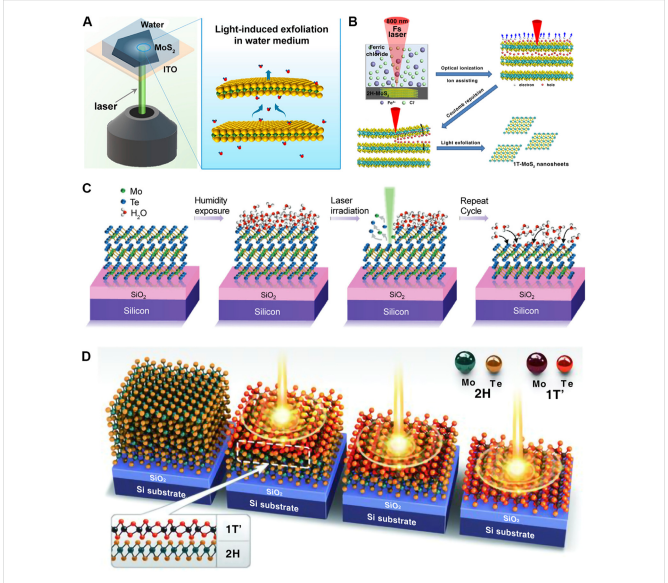


FIGURE 1
Top-down synthesis methods. (A) Schematic of the laser exfoliation device for 2H-MoS₂ flakes in aqueous media (Zhai et al., 2021) (Copyright 2021, American Chemical Society). (B) Schematic of the preparation of MoS₂ nanosheets through femtosecond laser exfoliation (Zuo et al., 2021) (Copyright 2021, American Chemical Society). (C) Schematic of the mechanism for laser thinning of MoTe₂ (Nagareddy et al., 2018) (Copyright 2018, Wiley-VCH). (D) Schematic of the laser-driven phase patterning process of MoTe₂ from 2H to 1T' (Cho et al., 2015) (Copyright 2015, American Association for the Advancement of Science).

2.2 Laser thinning

Laser thinning techniques have provided a direct and site-specific method for removing layers and obtaining "on-demand" 2D TMDCs (Castellanos Gomez et al., 2012; Lu et al., 2014; Sunamura et al., 2016; Kim et al., 2017; Park et al., 2017; Gong et al., 2018; Nagareddy et al., 2018; Rho et al., 2019; Tran-Khac et al., 2019; Wang et al., 2020; Kang et al., 2021). Unlike the traditional gas or plasma etching thinning method, laser thinning shows high accuracy on thinned layers, flexible programmable patterning mode, and high thinning efficiency. Nagareddy et al. (2018) demonstrated an efficient laser thinning method, which was constructed by the Raman spectroscopy and coupled with an atomic force microscope. The experimental setup was placed in

an airtight environment with strictly controlled humidity. Using ultra-low laser power, the thickness of MoTe $_2$ film can be controlled layer by layer from multilayer to monolayer, as shown in Figure 1C. In addition, the thickness reduction shows the linear correlation between the thickness reduction and the number of scans. The laser thinning process is achieved through the sublimation of the top layer, mainly caused by the conversion of light absorbed by the material into heat energy (Lu et al., 2014). The heat generated by the conversion of absorbed laser into energy is difficult to dissipate through the substrate due to the poor coupling between TMDC thin layers that are mediated by van der Waals forces (Su et al., 2021). Thus, until the laser power is increased, the bottom layer remains in close contact with the SiO $_2$ /Si substrate, acting as a heat sink to prevent removal.

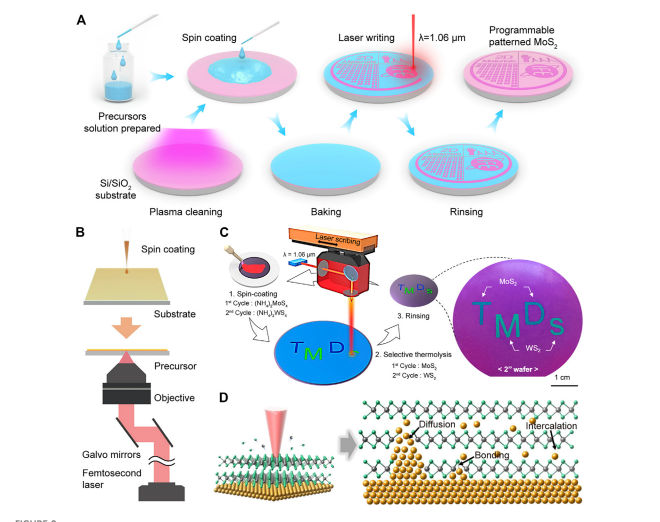


FIGURE 2
Bottom-up synthesis methods. (A) Flow diagram of direct laser writing, a method for the laser-directed synthesis of MoS_2 on the SiO_2/Si wafer (Xu et al., 2021a) (Copyright 2021, Elsevier). (B) Schematic of MoS_2 synthesized by femtosecond laser (Xu et al., 2022) (Copyright 2022, American Chemical Society). (C) Schematic of the layer synthesis of MoS_2-WS_2 heterostructure (Park et al., 2020a) (Copyright 2020, American Chemical Society). (D) Schematic of Au-doped MoS_2 (Huo et al., 2021) (Copyright 2021, American Chemical Society).

2.3 Laser driven phase

The exploration and manipulation of a novel phase of matter is a primary pursuit for materials research. The emergence of atomically 2D TMDCs has enabled the examination of diffusive, displacive, and quantum phase transitions. Cho et al. (2015) demonstrated a novel method of localized polymorph engineering and realized the laser-induced phase transition of MoTe₂. The multilayer 2H-MoTe₂ flake with about 30 layers was obtained by mechanical exfoliation. The schematic diagram of the phase transition mechanism is illustrated in Figure 1D. Under laser irradiation, the thickness of the irradiated region in 2H-MoTe₂ decreased by a few layers and transformed to 1T' phase. During the irradiation process, the 1T' MoTe₂ layer remained due to the heat sink effect of the SiO₂ substrate. It should be noticed that the phase transition is irreversible from 1T' to 2H phase even under the higher energy or intensity of the laser. The

driving force for the one-way phase transition is caused by the irreversible Te vacancy created under laser irradiation. Compared with conventional methods such as heat treatment, strain engineering, charge transfer, and plasma irradiation, laser-induced phase transition engineering allows for specific phase transition sites and patterned processing without impurities during the phase transition (Chen Z. R. et al., 2019).

3 Bottom-up synthesis methods

Unlike the top-down synthesis method, the bottom-up synthesis method is more efficient and cost-effective in the synthesis of 2D TMDC materials because it does not require the preparation of precursors using other methods. With the assistance of a laser, large-area patterned TMDCs, heterostructure, and atom doping can be realized.

3.1 Laser directed synthesis

Direct laser writing technology can directly pattern 2D TMDC materials without mask and lithography, reducing the process flow of device fabrication and the risk of chemical contamination. Generally, the fiber laser (1.06 μ m), carbon dioxide laser (10.6 µm), and femtosecond laser (780 nm) have been adopted for the synthesis of 2D TMDCs (Hu et al., 2018). Xu et al. (2021a) reported a method for the efficient synthesis of wafer-scale MoS₂ using a direct laser writing technique, and the workflow is shown in Figure 2A. With heating by a 1.06 µm commercial fiber laser, the MoS₂ can be programmable with text, lines, patterns, and matrices in a few minutes. The laser interacts with the precursor, and the reaction temperature facilitates the thermal decomposition of (NH₄)₂MoS₄ into MoS₂. Apart from the thermal decomposition, the photochemical reaction induced by two-photon absorption can also be adopted for synthesizing 2D TMDCs. As shown in Figure 2B, the femtosecond direct laser writing was carried out to synthesize MoS2 under an objective lens. Molybdenum acetylacetonate and carbon disulfide were adopted as the molybdenum and sulfur sources. The twophoton absorption occurs in the focal point, synthesizing MoS₂ from the molybdenum metal complex via a photochemical reaction. With the assistance of a femtosecond laser, a minimal line width of approximately 780 nm can be achieved.

3.2 Laser heterostructure

The direct laser writing technique allows the fabrication of single 2D TMDCs and the direct preparation of 2D TMDC heterostructures. Compared with the transfer method, the laser heterostructure process can be programmable patterned, avoiding material damage and pollution (Li et al., 2020). In addition, the synthesis efficiency of heterostructures is greatly improved compared with CVD. Park et al. (2020a) successfully prepared WS2-MoS2 heterostructures vertically by two-step laser scribing, as shown in Figure 2C. The MoS₂ layer was first synthesized by the thermal decomposition of (NH₄)₂MoS₄ when the temperature increased over 700°C under laser irradiation. While for the heterostructures, due to the different optical absorption coefficients, the MoS₂ layer shows a minor temperature increase, and only the (NH₄)₂WS₄ layer absorbed the laser energy and selectively decomposed into WS2. Thus, the selective growth process by laser effectively produces layer-by-layer 2D heterostructures in a programmable pattern.

3.3 Laser doping

The impurity atom can modulate the physical and chemical properties of semiconductors such as electrical and optical. A small amount of elemental doping could affect the energy bands enhancing semiconductor conductivity. However, doping elements in 2D TMDCs are relatively difficult and generate more defects in the 2D TMDCs. In addition, the weak

interaction will result in unstable interfaces due to the interaction between doped elements and 2D TMDCs (Mak et al., 2010). Laser provides a simple way for 2D TMDC doping, which is promising for future electronic applications (Chen J. et al., 2019). One way to achieve laser doping in 2D TMDCs is by dissolving the doping substance in the precursor, which results in doping during formation (Hu et al., 2020). In addition, doping of elements in preprepared 2D TMDCs can also be accomplished by introducing gas molecules or solids containing the desired doping atoms (Afaneh et al., 2018; Rho et al., 2019; Huo et al., 2021). For example, Huo et al. (2021) reported an Au-doped MoS2 process by doping a solid source, as shown in Figure 2D. MoS₂ was transferred to the Au electrode, and with femtosecond laser irradiation, the Au atom can be intercalated and diffused into MoS2. The interface diffusion and chemical bonding of Au reduced the Schottky barrier of the metal-semiconductor interfaces, which could enhance the performance of the devices.

4 Conclusion and outlook

This mini-review summarized the top-down and bottom-up methods for the laser synthesis of 2D TMDCs. Compared with the traditional methods, the laser synthesis methods have the advantages of flexibility, convenience, efficiency, and controllability. Laser synthesis technology can directly produce large-area, high-quality TMDCs, alloys, and heterostructure, which is a promising sign for future industrial-scale manufacturing.

Laser synthesis of 2D TMDCs is an emerging field, and some issues need to be further addressed, especially for the laser–reactive matter interaction. In addition, the synthesis of sulfides in the atmospheric environment is the focus of attention, while the atmosphere protection and hydrogen participation would be focused while synthesizing selenide and telluride. In addition, recent works on laser synthesis of 2D TMDCs constructed on the rigid substrate (such as SiO₂/Si) or liquid environments and direct laser synthesis on flexible substrates (such as polyimide, polydimethylsiloxane, polyethylene terephthalate, and polyethylene naphthalene) have not been realized yet, which would offer another chance for realizing applications in flexible electronics.

Author contributions

HW: investigation, methodology, visualization, and writing—original draft. MX: investigation, methodology, writing—original draft, writing—review and editing, visualization, and funding acquisition. HJ: investigation, writing—original draft, writing—review and editing, and visualization. TH: investigation, methodology, and conceptualization. WL: methodology, writing—review and editing, and funding acquisition. LZ: methodology, conceptualization, writing—review and editing, and funding acquisition. XW: conceptualization, methodology, writing—review and editing, supervision, project administration, and funding acquisition.

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Conflict of interest

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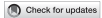
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Higher-order G-quadruplexes in promoters are untapped drug targets

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G-quadruplexes (G4s) are four-stranded nucleic acid secondary structures that form within quanine-rich regions of chromatin. G4 motifs are abundant in the genome, with a sizable proportion (~40%) existing within gene promoter regions. G4s are proven epigenetic features that decorate the promoter landscape as binding centers for transcription factors. Stabilizing or disrupting promoter G4s can directly influence adjacent gene transcription, making G4s attractive as indirect drug targets for hard-to-target proteins, particularly in cancer. However, no G4 ligands have progressed through clinical trials, mostly owing to off targeting effects. A major hurdle in G4 drug discovery is the lack of distinctiveness of the small monomeric G4 structures currently used as receptors. This mini review describes and contrasts monomeric and higherorder G-quadruplex structure and function and provides a rationale for switching focus to the higher-order forms as selective molecular targets. The human telomerase reverse transcriptase (hTERT) core promoter G-quadruplex is then used as a case study that highlights the potential for higher-order G4s as selective indirect inhibitors of hard-to-target proteins in cancer.

KEYWORDS

G-quadruplex (G4), higher-order G4, DNA, promoter, drug target

1 G-quadruplex structures in promoters: monomeric vs. higher-order forms

1.1 Intramolecular monomeric promoter G-quadruplexes

G-quadruplexes (G4s) are four-stranded secondary structures created from the stacking of two or more guanine tetrads ("G-tetrads") (Spiegel et al., 2020). Each G-tetrad is composed of four guanine bases arranged in a square planar configuration, that is, stabilized by Hoogsteen hydrogen bonding (Figure 1A). Monovalent cations are coordinated within the G-tetrad column central channel by the inward facing carbonyl groups, providing stabilization from coordinate bonding and neutralization of the partially negative charges (Lane et al., 2008). A commonly used sequence motif to describe a monomeric G-quadruplex is $G_{3-4}L_{1-7}G_{3-4}L_{1-7}G_{3-4}L_{1-7}G_{3-4}$, where G indicates a guanine tract and L designates any nucleotide in the intervening loop. Historically, the largest loop length has been taken to be seven nucleotides because of the destabilizing effect of large loops *in vitro* (Spiegel et al., 2020; Ravichandran et al., 2021).

G-rich sequences in specific gene promoter regions can adopt G4 structures with a surprisingly diverse array of topological conformations for the intramolecular monomeric G4 *in vitro*, depending on the sequence composition, loop lengths, ionic environment, and hydration (Chen and Yang, 2012; Miller et al., 2010; Bhattacharyya et al., 2016). The most

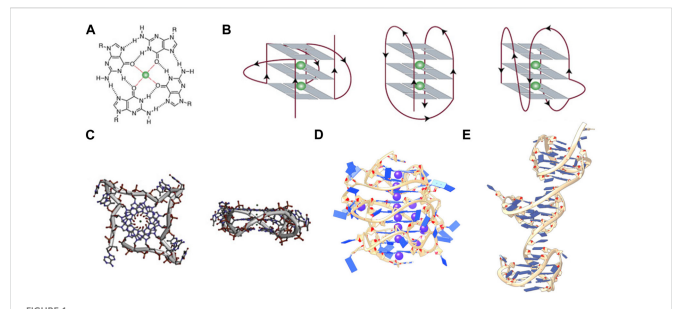


FIGURE 1
(A) Structure of a G-tetrad, (B) From left to right, schematic representations of (A) parallel, antiparallel, and mixed hybrid with bulge, (C) Top and side views of a parallel monomeric G-quadruplex, (D) side view of a stacked intramolecular all-parallel higher-order G-quadruplex, and (E) side view of an intramolecular higher-order all-hybrid G-quadruplex in a beads-on-a-string arrangement. (C–E) are not to scale. Figures 1A–C, are adapted from (Spiegel et al., 2020), (D) is adapted from (Monsen et al., 2021).

frequent topologies are: antiparallel, in which two out of four G-tracts run in the same direction; hybrid, in which three G-tracts run in the same direction; and parallel, in which all G-tracts run in the same direction (Figure 1B) (Spiegel et al., 2020). This review is primarily concerned with promoter G4s that are necessarily intramolecular and flanked on either end by single- or double-stranded regions. In the context of a promoter bubble (Shirude et al., 2007; Monsen et al., 2023), as well as when promoter G4 sequences are flanked by single-stranded regions (this can be realized in vitro, for example, by adding two residues on either end of the monomer G4 motif: L₂G_xL_nG_xL_nG_xL_nG_xL₂), the parallel topology is preferred (Chen et al., 2021). The biological preference for parallel promoter G4 conformations is supported by the extensive immunofluorescence seen across chromosomes stained with the parallel-favoring anti-G4 antibody "BG4" (Biffi et al., 2013; Javadekar et al., 2020). The first ever near-atomic structural study of a G4 in a near-native duplex bubble has shown that duplex end-stacking at the G-tetrad interface may be a key factor to promoter G4 stability, suggesting that antiparallel and hybrid topologies could be sterically prohibited (Monsen et al., 2023). Overall, in vitro monomeric promoter G4 sequences adopt a variety of topological configurations when isolated, but when placed in their biological context prefer the parallel form.

1.2 Intramolecular higher-order promoter G-quadruplexes

Intramolecular higher-order promoter G4s (xG4s) consist of two or more connected monomeric G4 domains. The xG4 motif is degenerate compared to the lower order G4 motif and is of the general form $(G_{2-4}L_{1-12}G_{2-4}L_{1-12}G_{2-4}L_{1-12}G_{2-4}L_{\ge 0})_n$ (Monsen et al.,

2022a), where the length of the 3' connecting loop can be zero and $n \ge 2$. The xG4 motif becomes substantially more degenerate when accounting for mismatches and/or bulges (i.e., non-guanine nucleotides within a putative G-tract) (Berselli et al., 2020). The xG4 motif is notable for multiple reasons. First, the zero-nucleotide connecting loop presents the possible situation where the last guanine of the 3' G-tetrad of the first G4 domain and the first guanine of the 5' G-tetrad of the second G4 domain create a continuous G-tetrad column that spans the entire macromolecule. If both G4s are parallel, this would facilitate formation of a single continuous medium groove spanning the two domains with a favorable locked 3' to 5' G-tetrad stacked interface (Kogut et al., 2019). A second, less obvious feature of the xG4 motif is that it accounts for two-tetrad G4s, in which there are only two guanines in the G-tracts. Two-tetrad monomeric G4s cannot exist as stable parallel conformations in vitro when isolated (Kejnovska et al., 2021), but have been seen in higherorder G4 assemblies (Monsen et al., 2022a) [we note that stable antiparallel two-tetrad G4s are seen in vitro (Lim et al., 2009)]. This is an important feature, as it reveals that biologically relevant xG4s have an expanded structural repertoire compared to the monomeric sequences. Lastly, xG4s seem to tolerate much longer loops (1-12+ nucleotides), both within and between the G4 domains (Berselli et al., 2020; Monsen et al., 2020; Monsen et al., 2022a). This may be the result of tertiary interactions that stabilize either across the G4-G4 domain by loop interactions (Rigo and Sissi, 2017), symmetry in the loop giving rise to a stabilizing hairpin moiety (Berselli et al., 2020), or arise from the stability imparted on the G4 cores from head-to-tail stacking (Kogut et al., 2019).

Structural characterizations of intramolecular promoter xG4s to date have revealed an overwhelming preference for stacked arrangements (Micheli et al., 2010; Rigo and Sissi, 2017; Monsen

et al., 2020; Monsen et al., 2022a), with the highest resolution models showing a preference for an all-parallel stacked globular configuration (Figure 1D) (Monsen et al., 2020; Monsen et al., 2022a). Importantly, all reported putative xG4 sequences that have been investigated form stable G4 structures in vitro. In some cases, the presence of long loops (>7 nucleotides) result in hairpins (Monsen et al., 2022a) that contribute to the stability of the individual domains (Ravichandran et al., 2021) while also creating structurally unique interfaces. In contrast to promoter xG4s, other higher-order intramolecular G4s reported, such as the human telomere (Monsen et al., 2021), the insulin-linked polymorphic region (ILPR) minisatellite (Schonhoft et al., 2009), and the CEB25 minisatellite locus (Amrane et al., 2012), show mixed antiparallel and hybrid topologies that are better described as dumbbell shaped or beads-on-a-string configurations. Figure 1 contrasts the structures of a monomer G4 in the parallel conformation, a higher-order all-parallel stacked G4, and a higher-order all-hybrid beads-on-a-string.

The selective binding interface of the xG4s is not imparted by the G-tetrad columns, but rather the specific G4-G4 interaction interfaces, loop sequence, size, and configurations. Figure 1C reveals that there is little targetable real estate associated with the monomeric parallel G4 (Monsen et al., 2023). Conversely, Figures 1D, E reveals multiple putative binding sites that would be large enough and distinct enough for specific protein or drug interaction. Biochemical support for this idea comes from a recent pull-down study conducted by Ceschi et al. (Ceschi et al., 2022). In this study, the authors used a variety of higher-order G4 sequences (such as those in Figures 1D, E) to enrich for tightly interacting proteins from cell lysates that are specific to higherorder G4s over the lower order forms. Surprisingly, the intermediate filament Vimentin was shown to have a selective nanomolar affinity to higher-order G4 structures with no apparent binding to lower order species. While the mechanism of recognition is still unclear, this study supports the hypothesis that xG4s offer unique recognition sites that could be useful in selective targeting.

2 G-quadruplex distribution and function in promoters

2.1 Biological distribution and function of promoter G4s

Promoter G-quadruplexes are prevalent epigenetic regulatory elements. Current estimates show more than 700,000 monomeric G4 motifs across the human genome (Hansel-Hertsch et al., 2017). Huppert and Balasaubramanian, using the canonical sequence motif $G_{3+}L_{1-7}G_{3+}L_{1-7}G_{3+}L_{1-7}G_{3+}$, showed that more than 40% of gene promoters have at least one monomeric G4 motif (Huppert and Balasubramanian, 2007). More recently, Hänsel-Hertsch and colleagues have used a chromatin immunoprecipitation (ChIP)-sequencing direct capture approach to show that thousands of G4 structures, not just motifs, are enriched in highly transcribed gene promoters, specifically many involved in cancer (Hansel-Hertsch et al., 2016). Promoter quadruplexes overall appear to be acting as general transcription factor (TF) "binding hubs",

coinciding with regions of open chromatin and high transcriptional activity (Spiegel et al., 2021).

At a more granular level, promoter G4s act in concert with transcriptional proteins to affect gene transcription in multiple ways. Initially, promoter G4s were thought to only act as simple physical barriers to polymerases, acting as "on/off" switches of transcription (Sarkies et al., 2010). However, studies have now shown that G4s can directly recruit transcription factors with some level of specificity. For instance, two zinc fingers, SP1 and MAZ (Myc-associated zinc finger), show G4 structure-dependent recognition. In the former case, Raiber and colleagues, using pull-down experiments with the transcription factor SP1, showed that 36% of the sequences lacked consensus SP1 binding motifs (Raiber et al., 2012). They went on to show that 77% of those sequences lacking the SP1 consensus motif were putative G-quadruplexes and that, overall, SP1 binding had 87% overlap with G4 sequence motifs. In the latter case, Cogoi and colleagues have shown that MAZ recognizes a G4 formed within the kRas promoter and showed that stabilizing the kRas G4 with a small molecule could promote MAZ binding and increase transcription, while mutations that destabilized the G4 reduced MAZ binding and transcription (Cogoi et al., 2010).

G4s can also serve as transcriptional repressors. The classical case for this is the G4 formed within the *c-Myc* promoter. In their seminal study, Siddiqui-Jain et al. showed that a G-rich region in the nuclease hypersensitivity element III (NHEIII) of the c-Myc P1 promoter forms one or more monomeric G4s (Siddiqui-Jain et al., 2002). Through mutational destabilization, they showed that there is a 3-fold increase in c-Myc expression, indicating that the G4 acts as a transcriptional repressor. They showed that stabilization of the G4 with a small molecule could further reduce transcription to below the basal level. Later studies on the protein nucleolin, a multifaceted and abundant protein found in the nucleolus (Tajrishi et al., 2011), have shown that it can help the folding of promoter G4s like a molecular chaperone (Tosoni et al., 2015). Nucleolin was shown to fold the c-Myc NHEIII G4, promoting transcriptional downregulation in cells (Gonzalez and Hurley, 2010). Altogether, these studies show that promoter G4s are important epigenetic regulators of genes, and that their stability and interaction with transcription factors can influence transcription.

2.2 Biological distribution and function of promoter xG4s

Promoter xG4s are also abundant across the genome. Berselli and colleagues recently developed QPARSE (13), the first algorithm capable of finding xG4s in the genome that accounts for mismatches and bulges in G-tracts (see Figure 1B for a G4 with a bulge). In the study, the authors used their degenerate G4 motif algorithm to find monomeric, dimeric, and trimeric G4 sequences in the range of –200 to +600 of the transcriptional start sites (TSS) across the annotated human genome [GENCODE (Harrow et al., 2012)]. They found that 49%, 15%, and 4% of TSS regions had monomer, dimer, and trimer G4 repeats, respectively, and show that this enrichment cannot entirely be attributed to high GC content. Ceschi et al. have recently used the same algorithm to search just the first 100 bp upstream of the TSS of gene promoters in GENCODE, finding

1,478 dimer and trimer promoter xG4s (Ceschi et al., 2022) (\sim 4% of the 38,404 annotated genes in GENCODE v34).

Parsing out the *in vivo* function of a particular xG4 compared to their monomeric counterparts is a challenging task. At one end you have bioinformatic and G4-or G4-ChIP-sequencing approaches that lack the spatial resolution to distinguish between monomer, dimer, trimer, and other higher-order G4 effects (Park, 2009; Mahony and Pugh, 2015). For instance, the ChIP-sequencing studies mentioned above should, in theory, encompass promoter sites enriched with xG4s. At the other end, there is a dearth of tools that allow direct probing of the effects of monomer versus higher-order G4s. Specifically, there are no reports to date that have convincingly shown selective (de)stabilization of a single G4 domain among a higher-order promoter G4 assembly (Frasson et al., 2022). Mutational reporter assays are the current best approach to parsing out the functionality of xG4s within a cellular context (Siddiqui-Jain et al., 2002; Cogoi et al., 2010), but these come at the risk of altering protein recognition motifs (Bell et al., 2015).

The most thorough investigation of a promoter xG4 to date was conducted in 2019 by Ducani and colleagues on an xG4 found in the promoter of the proto-oncogene c-Kit (Ducani et al., 2019). The c-Kit gene encodes for a transmembrane tyrosine kinase receptor (c-Kit or CD117) that, after activation by stem cell factor (SCF), transduces signals that promote cell proliferation, differentiation, and migration (Liang et al., 2013). Excessive signaling by continued stimulation or mutation has been implicated in a variety of cancers, such as gastrointestinal stromal tumors (GISTs), pancreatic cancer, melanoma, and hematological neoplastic diseases (Gregory-Bryson et al., 2010; Abbaspour Babaei et al., 2016). Since c-Kit has a tyrosine kinase domain, these cancers are typically treated with tyrosine kinase inhibitors (TKIs) tailored to the mutation type (Abbaspour Babaei et al., 2016). Unfortunately, treating c-Kit-driven cancers with TKIs is often followed with a rapid switch to drug-resistance through mutation (Demetri et al., 2002; Loughrey et al., 2006; Abbaspour Babaei et al., 2016). Therefore, understanding how the *c-Kit* promoter xG4 regulates its transcription is important for drug development efforts. The c-Kit xG4 consists of three monomeric G4 motifs, designated K2, SP, and K1 (from 5' to 3') connected by a single dA loop and a hexanucleotide dGCGCAG loop, respectively. Each separate G4 domain has been structurally and/or functionally examined (Rankin et al., 2005; Phan et al., 2007; Hsu et al., 2009; Kuryavyi et al., 2010; Raiber et al., 2012; Kotar et al., 2019), and the higher-order structural assembly of domains K2-SP confirmed by integrative structural biology approaches (Rigo and Sissi, 2017; Monsen et al., 2022a). Both structural studies show that the K2-SP regions interact through stacking, although the two studies differ slightly in the size of the sequence examined. Based on circular dichroism of the full-length sequence used by Ducani, however, the full-length K2-SP-K1 sequence adopts an all-parallel conformation, which is consistent with the promoter xG4s structurally verified to date (Monsen et al., 2020; Monsen et al., 2022a). To investigate the biological function of such an arrangement, Ducani and colleagues conducted luciferase assays to test the effect of disrupting all combinations of the G4 regions by mutating guanines essential for structural integrity. They show in the leukemia cell line HEL92.1.7 that the K1 G4 (alone) and K2-SP G4s (together as a higher-order feature) have opposing roles in transcriptional regulation, with the former acting repressively and the latter acting to stimulate transcription. Further, they show that each G4 unit is significantly affected by the formation or disruption of the others, signifying

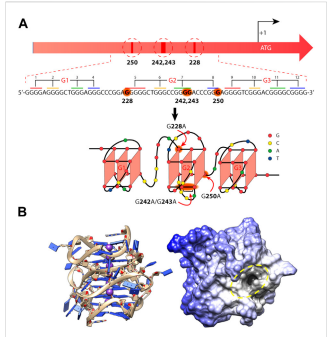


FIGURE 2 Mutations and folding of the hTERT core promoter G-quadruplex. (A) Cancer-specific mutations in the hTERT promoter's primary structure that generate de novo binding sites for the ETS transcription factors and putative structure that was proposed for the G-rich (noncoding) strand of the hTERT promoter. Below is the sequence of the WT hTERT G4 motif and one conceivable way of G4 folding. The assignment of G-tracts (numbered) to three G4motifs (G1, G2, and G3) is presented. Common mutation sites are shown in red in the hTERT G-rich sequence and in red arrows in the structural model. (B) The folded all-parallel hTERT xG4 represented with sugar-phosphate backbone as ribbons and nucleobases as slabs (left) and surface representation showing a large binding pocket (dark area inside of yellow dashed oval) at the junction between the first and second G4s. (A) was adapted from (Pavlova et al., 2022) and (B) was adapted from (Monsen and Trent, 2018)

G4-G4 crosstalk within a cellular context. The readout from each mutational state was more of a continuous distribution, rather than an all or nothing response. Overall, this study suggests that xG4s play a very nuanced role in governing transcriptional activation or repression. It remains to be seen whether a K2-SP disrupting small molecule, or SP-K1 stabilizing small molecule will act as an indirect c-Kit inhibitor.

3 Case study: targeting the hTERT core promoter xG4

One of the most studied xG4s to date is found within the *hTERT* core promoter region. *hTERT* encodes the protein catalytic subunit of telomerase, the ribonucleoprotein primarily responsible for maintaining telomere length homeostasis (Bryan and Cech, 1999). Although typically undetectable in somatic cells, hTERT is aberrantly overexpressed in more than 90% of aggressive cancers (Shay and Bacchetti, 1997), making it a long sought after cancer-specific target. The *hTERT* gene was first identified as harboring a putative xG4 in its promoter by Palumbo et al. (2009). The hTERT xG4 sequence consists of twelve G-tracts of three or more guanines that enable the maximum formation of three contiguous G4s (Figure 2).

Since its discovery, the tertiary structure of the hTERT xG4 has been intensely disputed. In the original work, Palumbo et al. showed by CD that the hTERT sequence was entirely parallel (Palumbo et al., 2009), although, based on their DMS footprinting results, they proposed a model consisting of a parallel G4 unit connected to an antiparallel G4 unit connected by a ~26 nucleotide hairpin loop. About a year after, Micheli et al. (2010) independently proposed that the hTERT xG4 formed three parallel G4 units that were contiguous and stacked 3' to 5' based on CD and Taq polymerase stop assays. Later studies using an experimental small molecule and DMS footprinting assays reported yet another, slightly different, G4-hairpin arrangement (Song et al., 2019). However, structural studies of the sequence using a combination of CD, hydrodynamics, NMR, modeling, and small-angle X-ray scattering (SAXS) have since confirmed that the most consistent model is one in which three parallel G4s are tightly stacked in a stacked 3' to 5' arrangement (Figure 2) (Monsen et al., 2020). A recent DMS footprinting experiment confirms the all-parallel model as correct (Pavlova et al., 2022).

The hTERT xG4 is a potentially selective, transcriptionally repressive structure. The hTERT core promoter region has multiple non-coding mutations that are seen across many cancer types, with the two most prominent denoted "G228A" and "G250A", that coincide with a robust increased telomerase activity (Killela et al., 2013). These mutations exist within G-tracts that form the central G4 unit of the three stacked all-parallel xG4 model (see Figure 2) (Micheli et al., 2010; Chaires et al., 2014; Monsen et al., 2020). Using luciferase expression assays of the wild type (WT) promoter versus mutant promoters of either G228A or G250A, Bell and colleagues showed that a robust increase in promoter activity is gained with either mutant (Bell et al., 2015). Further investigation showed that either G- > A mutation creates a de novo ETS (Erythroblast Transformation Specific) transcription factor consensus motif that is recognized by the transcription factor GABP (GA-binding protein). However, the creation of an ETS motif has been shown to be insufficient to fully explain the transcriptional changes seen in mutant cells (Kang et al., 2016).

Early investigations by Micheli et al. showed that the central G4, which encompasses either G- > A mutation, is unstable as an isolated monomeric G4, and only forms stably as a higher-order G4 assembly through stacking with both 5' and 3' G4 regions (Micheli et al., 2010). The destabilizing effects of the mutants on the overall xG4 have since been confirmed (Pavlova et al., 2022). A G4 ligand reported to refold and stabilize the hTERT xG4 WT structure (with or without G- > A mutations) was able to restore its repressive effects and showed good selectivity for the hTERT promoter over other genes with known promoter G4s (Kang et al., 2016). More recently, Monsen et al., (2022b) used a virtual screening approach to target the loop and G4-junctional regions of the all-parallel hTERT xG4 (Figure 2). Using a variety of in vitro binding and competition assays, the authors were able to find a drug-like small molecule that stabilized across the second and third G4 regions that showed high selectivity over duplex DNA and moderate selectivity over all other forms of DNA tested. In both cases, repression of hTERT expression was confirmed in breast cancer cell lines. Collectively, these studies reveal that the hTERT xG4 is an indirect target for down-regulating hTERT in cells. Further, these studies show for the first time that a higher-order promoter G4 can be targeted with selectivity using the unique features imparted from G4-G4 domains.

4 Discussion

Higher-order G-quadruplexes have emerged as selective targets in the promoters of thousands of annotated genes across the human genome. While xG4s offer some of the same protein binding recognition that monomeric G4s do, promoter xG4s also appear to encode for transcriptional status, possibly acting as titratable "dimmer switches" of gene activity. Structurally, xG4s offer a much richer drug targeting landscape consisting of G4-G4 interaction junctions adorned with protein-like binding pockets formed among sequence-specific loop features. This is emphasized by the decadelong pursuit targeting the *hTERT* core promoter xG4 which has resulted in both a unique receptor and the first drug-like small molecule targeting an xG4 with selectivity.

While xG4 studies to date have led to exciting new insight into their structural arrangements and potential regulatory mechanisms, there is still much to be understood. One of the major hurdles in studying xG4s is their recalcitrance to the traditional structural biology techniques NMR and X-ray diffraction. To date, all structural models have been derived from medium-to lowresolution integrative structural strategies and/or footprinting methods (Palumbo et al., 2009; Micheli et al., 2010; Rigo and Sissi, 2017; Monsen et al., 2020; Monsen et al., 2021; Monsen et al., 2022a; Pavlova et al., 2022). Cryo-EM is emerging as a possible solution to this problem (Monsen et al., 2023); however, atomic resolution of relatively small, potentially heterogeneous [e.g., G-tract isomers (Harkness and Mittermaier, 2017; Hennecker et al., 2022)] or flexible systems (e.g., inter-domain movements or long flexible loops) (Monsen et al., 2021; Monsen et al., 2022a; Monsen et al., 2023) is still a considerable challenge in the field (Herzik et al., 2019) [although scaffolds might offer a solution (Wu and Rapoport, 2021)]. A second major hurdle, as touched on above, is determining their biological mechanism. Molecular tools, such as xG4-specific fluorescent molecules (Summers et al., 2021) or antibodies (Biffi et al., 2013), should aid in revealing their spatial and temporal formation. Recently, the zinc finger-containing transcription factor Yin Yang-1 (YY1) was shown to bind G4s and bring two G4 domains into proximity through its dimerization (Li et al., 2021), offering a potential tool for investigating biological function of xG4s in a more biologically relevant context.

Author contributions

RM conceptualized the topic, researched, and analyzed the literature, and wrote the manuscript.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Transition metal anchored on red phosphorus to enable efficient photocatalytic H₂ generation

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Transition metal (TM) single atom catalysts (SACs) are of great potential for photocatalytic H₂ production because of their abundant catalytic active sites and cost-effectiveness. As a promising support material, red phosphorus (RP) based SACs are still rarely investigated. In this work, we have carried out systematic theoretical investigations by anchoring TM atoms (Fe, Co, Ni, Cu) on RP for efficient photocatalytic H2 generation. Our density functional theory (DFT) calculations have revealed that 3d orbitals of TM locate close to the Fermi level to guarantee efficient electron transfer for photocatalytic performances. Compared with pristine RP, the introduction of single atom TM on the surface exhibit narrowed bandgaps, resulting in easier spatial separation for photongenerated charge carriers and an extended photocatalytic absorption window to the NIR range. Meanwhile, the H₂O adsorptions are also highly preferred on the TM single atoms with strong electron exchange, which benefits the subsequent water-dissociation process. Due to the optimized electronic structure, the activation energy barrier of water-splitting has been remarkably reduced in RPbased SACs, revealing their promising potential for high-efficiency H₂ production. Our comprehensive explorations and screening of novel RP-based SACs will offer a good reference for further designing novel photocatalysts for high-efficiency H_2 generation.

KEYWORDS

red phosphorus, single-atom catalysts, transition metals, photocatalysis, H₂ generation

1 Introduction

Hydrogen energy is a clean and renewable energy, attracting increasing research attention in recent years. Especially, it has a high gravimetric energy density, and the product of H₂ combustion is only pure water, making it a promising candidate to cope with the fossil energy crisis and meet the challenge of peak carbon dioxide emission limitation. (Wang S. et al., 2019; Pan et al., 2021; Xue et al., 2022). However, the high cost of H₂ production as well as safety issues in storage and transportation have greatly restricted its commercial applications. Without additional electric power consumption, solar energy-driven catalytic water splitting has been considered as an eco-friendly strategy for low-cost H₂ production. Currently, photocatalytic H₂ evolution has not been broadly adopted in large-scale industrial production since it still suffers from the low solar-to-hydrogen (STH) conversion rate, which is caused by intrinsic unfavorable thermodynamics and sluggish kinetics of solar-driven catalytic water splitting reaction. (Chen et al., 2020; Bie et al., 2022). In recent decades, to improve the catalytic activity of photocatalysts, researchers have tried a

series of modulation strategies to enhance surface reactivity, reduce the activation energy barrier, and restrain the intrinsic fast recombination of photoinduced charge carriers.

Previously, phosphorus has been treated as an effective heteroatom dopant for the band structure engineering of TiO2based photocatalysts. (Yu et al., 2003; Wang et al., 2012a; Hu et al., 2017; Zhu et al., 2019; Huang et al., 2022). Afterwards, it has been discovered that phosphorus itself can also act as an efficient photocatalyst for hydrogen evolution through water-splitting. (Shen et al., 2014; Hu et al., 2017; Liu et al., 2019; Chen et al., 2022). Among its three allotropes (black, P₄-white, and red phosphorus), red phosphorus (RP) is the most available form and has excellent chemical stability at room temperature, and it has been treated as a promising semiconductor material in the construction of photocatalytic platforms. (Wang et al., 2012a; Xia et al., 2015; Hu et al., 2017; Liu et al., 2019; Chen et al., 2022; Fung et al., 2022). It has been reported that RP is a durable photocatalyst with the stable catalytic activity of H2 evolution for more than 90 h under visible light irradiation. (Wang et al., 2012a; Wang et al., 2012b). Though RP has good absorption of visible light at about 700 nm, the overall water-splitting efficiency of pristine RP is relatively limited due to its low conductivity, poor electron-hole separation efficiency, and sluggish charge-carrier mobility. (Zhu et al., 2019; Wang M. et al., 2020; Wu et al., 2021; Chen et al., 2022).

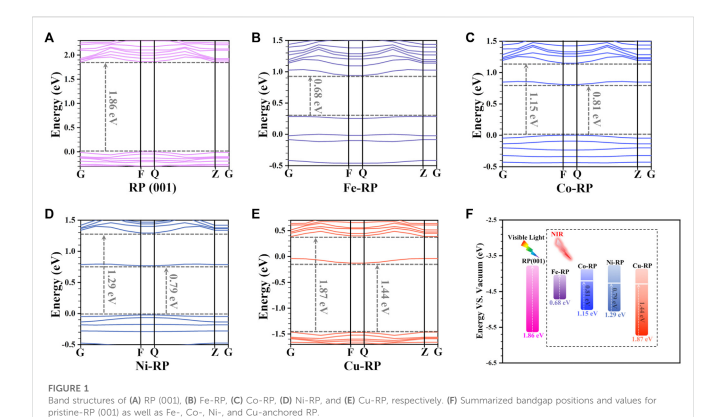
Since RP has good lattice compatibility, it is possible to modify the catalytic surface through the doping method or heterostructure construction. As a result, the electronic structures of RP can be wellmanipulated to benefit the photocatalytic reaction. (Li et al., 2019; Zhu et al., 2020a; Huang et al., 2022; Wang et al., 2022; Jia et al., 2023). With the assistance of appropriate active co-catalyst atoms such as Pt, the overall photocatalytic H₂ production performance of RP is greatly improved. (Wang et al., 2012a). In consideration of material cost and future industrial applications, developing nonprecious metals based catalysts is of great significance for the popularization of hydrogen energy. In particular, 3d transition metals (TMs) are the most commonly studied candidates to replace noble-metal catalysts due to their high catalytic activity. By suitable tuning strategies, the novel non-noble metal catalysts can also achieve comparable or even superior catalytic activity to noble metal catalysts. Scientists have reported that both morphological and scale controls are effective for the electronic property tuning of semiconductor materials. (Zhu et al., 2020b; Wu et al., 2020; Khandelwal et al., 2022). By downsizing the material scale from the nanoscale to the cluster scale and then finally the atomic scale, an enhanced inter-atomic strain is induced, leading to the modulations in geometrical structure and electronic structures. (Liu and Corma, 2018; Khandelwal et al., 2022). Hence, developing single-atom catalysts (SACs) is a potential method for the modulation of catalytic activity on the atomic scale. The atomic dispersion of active metal atoms on the RP surface will maximumly improve the atomic utilization rate, offering maximum photocatalytic active sites to enable the light adsorption and activation of H₂O molecules with a minimum consumption of metal atoms. (Wang B. et al., 2019; Gao et al., 2020). In previous studies, the Ni-anchored RP SACs have been proven to be very efficient photocatalysts for water splitting, with a greatly improved catalytic activity than the pristine RP. (Wang et al., 2022; Jia et al., 2023). However, the detailed

photocatalytic reaction mechanisms of Ni-RP SACs and the potential of other 3d TMs SACs still lack sufficient studies.

In this work, we have conducted comprehensive calculations of the electronic structures of commonly used TMs-based SACs (TM = Fe, Co, Ni, and Cu) on RP support (denote as TMs-RP) to predict their performances of photocatalytic H₂ generation. Based on the band structures and work functions, the thermodynamic driving force for hydrogen and oxygen evolution reaction (HER and OER) of each TMs-RP SACs is obtained and the pH tuning is also realized based on the Nernst equations. Besides, the kinetics for water adsorption and activation, the energy barrier for H adsorption, and the activation barrier of H₂ evolution are demonstrated in detail. This work will offer an atomic scale insight into the reaction mechanisms of TMs-RP SACs based photocatalytic H₂ evolution. Meanwhile, the detailed theoretical explorations also serve as fundamental theoretical references for the rational design of TMs-based SACs with enhanced photocatalytic performances in the future.

2 Calculation setup

In this work, we have conducted density functional theory (DFT) calculations within the CASTEP module. (Clark et al., 2005). The geometry optimizations and single-point energy calculations for all models have been conducted based on the GGA-PBE (Perdew et al., 1992; Perdew et al., 1996; Hasnip and Pickard, 2006) functional, ultrasoft pseudopotentials, and the BFGS algorithm. To guarantee the electronic minimization and convergence requirement, we have adopted the ensemble DFT method of Marzari et al. for the solution of the Kohn-Sham equation. (Marzari et al., 1997). The cutoff energy has been applied with the ultrafine quality, which is set as 380 eV for RP and Fe-RP, Co-RP, Ni-RP, and 440 eV for Cu-RP. The k-point set is $2 \times 2 \times 1$ for all the energy minimizations. The convergence parameters are set up as follows: the SCF tolerance is 5.0 \times 10⁻⁷ eV/atom; the Max. Hellmann-Feynman force per atom is 0.01 eV/Å; the Max. stress is 0.02 GPa; and the Max. displacement is $5.0 \times 10^{-4} \,\text{Å}$. According to previous experimental studies, the [001]-oriented Hittorf's phosphorus has been proven to be very efficient for water-splitting. (Zhu et al., 2020a). Thus, in this work, we choose the RP (001) surface to act as the supporting host to stabilize the single-atom (SA) TMs. The RP (001) facet is built from relaxed bulk Hittorf's phosphorus, (Thurn and Krebs, 1966), with a thickness of two layers including 84 atoms (Supplementary Figure S1A). The top view of pristine RP (001) and TMs-RP (Fe, Co, Ni, and Cu) catalysts are demonstrated in Supplementary Figures S1B-F. The lattice parameter is 9.27 Å and 9.21 Å for length A and B-orientations, respectively. The vacuum thickness is set to be 20 Å, resulting in a length of 40.95 Å in C-orientation to ensure sufficient space for geometry optimizations. During the adsorption of H₂O and H on different TMs-RP in this work, the SAC surfaces have been constrained in order to highlight the behaviors of key adsorbates. The H-adsorption free energy (ΔG_{H^*}) serves as a key descriptor for the prediction of HER activity, which can be calculated based on the following equation (Nørskov et al., 2005; Kerketta et al., 2022):



$$\Delta E_{\rm H} = E_{\rm H^*} - E^* - 1/2E(H_2) \tag{1}$$

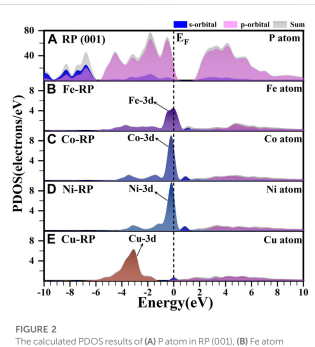
In this equation, * indicates the un-adsorbed pure surface, and H* stands for the surface adsorbed with H.

3 Results and discussions

3.1 Electronic structures of TMs-RP

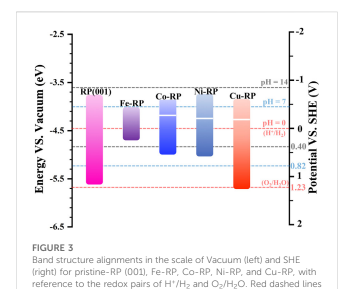
For photocatalytic reactions, the excitation energy required for electron-hole separation is largely decided by the bandgap value of the semiconductor photocatalyst. To make the most use of solar energy, the bandgap of the photocatalysts is expected to be located within the spectrum of sunlight, namely, from the UV to NIR range. In fact, most of the solar energy (nearly 95%) is constituted by visible light (43%) and NIR (52%). To match the Vis-NIR excitation window of sunlight, the bandgap of the candidate photocatalyst is expected to be less than 3.10 eV. And based on the redox potential of water, to drive the water splitting reaction, the bandgap value is projected to be wider than 1.23 eV with the location of VBM lower than the oxidation level of O2/H2O and CBM higher than the reduction level of H₂/H⁺. Thus, the bandgap values and band positions of photocatalysts can largely reflect their absorption window toward sunlight, and the relative positions of VBM and CBM with reference to standard hydrogen electrode potential can be used to predict the redox tendency for water splitting.

To evaluate the bandgap matching degree of the TMs-RP, we have first compared the band structure of pristine RP (Figure 1A)



The calculated PDOS results of **(A)** P atom in RP (001), **(B)** Fe atom in Fe-RP **(C)** Co atom in Co-RP, **(D)** Ni atom in Ni-RP, and **(E)** Cu atom in Cu-RP respectively. The dashed line means the Fermi level.

with Fe-, Co-, Ni-, Cu-anchored RP, respectively (Figures 1B–E) to reveal the anchoring effect from different TM atoms. For the pristine RP (001) surface, the bandgap is calculated to be 1.86 eV,



indicate pH = 0, blue dashed lines represent pH = 7, and black

dashed lines stand for pH = 14.

corresponding to an excitation energy of 667 nm in the red-light range. For Fe-, Co-, and Ni-anchored RP, we find additional TM states generated upper to the valence band maximum (VBM) and below the conduction band minimum (CBM), resulting in a narrowed bandgap of 0.68 eV, 1.15 eV, and 1.29 eV, respectively. Notably, the bandgap of Cu-RP is not affected by Cu single atoms, which still remains 1.87 eV. In particular, owing to the states appeared below CBM in Co-RP (0.81 eV) and Ni-RP (0.79 eV), the excitation energy required for the generation of photon-induced charge carriers has largely shifted to the NIR range (1531 nm–1824 nm), as indicated in Figure 1F. For Cu-RP, there

is an obvious downshifting of both VB and CB, which shows only

one mid-gap state with a distance of 1.44 eV from the VBM, which

reduces the absorption energy to a lower NIR window (861 nm).

Besides band structures, the calculated partial density of states (PDOS) is another important indicator of electronic structures. In Figure 2, we focus on the analysis of the d-orbital property of each TM since the d-band center is also strongly related to the photocatalytic activity and the electron transfer efficiency. (Zhou et al., 2022). The pristine RP shows an evident bandgap between CBM and VBM, which are both dominated by the P-3p orbitals (Figure 2A). For Fe-, Co-, and Ni-anchored RP, the localized d orbitals locate very close to the Fermi level (E_F) and VBM (Figures 2B-D). Especially, Co-RP and Ni-RP both display very sharp 3d orbitals with higher electron density near E_F, exhibiting better performance for electron transmission. In contrast, the 3d orbitals of Cu-RP have significantly moved to a lower position of VB, which is far from the E_F (Figure 2E). This results in a wider gap for the transition of photo-induced charge carriers during photocatalysis. The band edge is mainly contributed by the p-orbitals of P with overlapping of 3d orbitals of TMs, which induces the p-d coupling effect to further facilitate charge-carrier transfer. For all four TMs-RP catalysts, we notice the contribution of s-orbital to the CBM, which serve as a ladder to lower the excitation energy for electron excitation from VB to CB. For Cu-RP, the s-orbital dominates the $E_{\rm F}$, which only limitedly compensates for low electron transfer density near the VBM. Overall, the PDOS results have revealed that the single-atom TM anchored in RP, especially Co- and Ni-, brings great improvements in both separations and transportations of photo-generated charge carriers during photocatalysis.

3.2 Band structure alignment analysis

The electron-transition capacity of valence electrons is decided by the energy gap between E_F and vacuum level. (Zhou et al., 2022). Thus, the work function has been treated as a key descriptor for electron transition and charge flow in heterojunction structure, and it can be obtained from the calculated electrostatic potential. (Qin et al., 2020; Ruan et al., 2022). As discussed above, besides the bandgap values, the relative position of the bandgap with a reference to standard hydrogen electrode potential is also critical to predict photocatalytic activity. For the construction of the band alignment diagram (Figure 3), the electrostatic potential for each TM-RP type studied in this work has been calculated to indicate the distance between the Fermi level and vacuum level (Supplementary Figures 2A-E). The thermodynamic driving force for photocatalysis is mainly affected by the relative potential position of CBM and VBM of the semiconductor as well as the redox potential of the corresponding reversible reaction. (Li et al., 2016). More negative CBM means stronger reduction reaction tendency, while more positive VBM potential signifies stronger oxidation reaction driving force. Therefore, besides a bandgap value of more than 1.23 eV, the band edges (both VBM and CBM) of the photocatalyst candidates need to cover both the standard potentials of OER and HER to guarantee the high performances of water-splitting. (Mortazavi et al., 2021).

In the acidic condition (pH = 0 in aqueous solution), the CBM of pristine RP and TM-anchored RP are all located above the reduction level of H₂/H⁺, exhibiting a high hydrogen evolution tendency (Supplementary Figure S3). However, based on the energy gap between CBM and H₂/H⁺ level, the TMs-RP all exhibit a relatively weaker hydrogen evolution tendency than the pristine RP surface except Ni-RP. According to the large gap value from Ni-RP CBM to the H₂/H⁺ level, we propose that the anchoring Ni atom offers a stronger driving force for the hydrogen evolution reaction. Meanwhile, the VBM positions of pristine RP and TM-RP (except Cu-RP) are located above O₂/H₂O level, theoretically indicating the weak capability for oxygen evolution. Previous works have revealed the slight mismatching of bandgaps and work functions between theoretical calculations and experimental data. (Wang et al., 2009; Wang et al., 2012a). By considering the calculated SHE as a reference, the pristine RP and Cu-RP are also able to achieve oxygen evolution in the acidic condition.

Since standard redox potentials of H₂O are sensitive to pH variation, the photocatalytic driving force for HER and OER can be well tuned by the pH manipulation based on the following Nernst equations: (Shojaei et al., 2020; Mortazavi et al., 2021):

$$E^{HER}(H^+/H_2) = -4.44 \,\text{eV} + \text{pH} \times 0.059 \,\text{eV}$$
 (2)

$$E^{OER}(O_2/H_2O) = -5.67 \,\text{eV} + \text{pH} \times 0.059 \,\text{eV}$$
 (3)

The pH variation also impacts the band positions, which has been verified in experiments that the slope of E_{CB} (-0.033 eV/pH) does not follow the Nernstian dependence on pH (-0.059 eV/pH). (Simon et al., 2014). The slope of Nernstian dependence on pH for H₂O redox potentials is steeper. However, in this work, the pH value influences on the corresponding band positions are not included. In our work, the model is built in a vacuum environment with the mainly related elements (H2O, H, or OH) adsorbed on the catalytic surface in HER. The corresponding band positions are stationary with redox potentials of H₂O in different pH values to reveal the reaction tendency for HER and OER. (Zhang et al., 2021; Yang et al., 2018; Makaremi et al., 2018; da Silva et al., 2019; Shahid et al., 2020). In the neutral condition (pH = 7), the H_2/H^+ level and O_2/H_2O level move up to a more negative position (with reference to SHE) at -0.41 eV and 0.82 eV, respectively (Supplementary Figure S4). Accordingly, for the pristine RP and TMs-RP photocatalysts, the driving force for HER declines while the tendency for OER increases. RP and Cu-RP still cover both potential levels of HER and OER, indicating their photocatalytic activity towards the full water splitting under neutral conditions. In addition, in the highly alkaline environment (pH = 14), pristine RP and TMs-RP nearly lose their HER capability in terms of thermodynamics (Supplementary Figure S5). Thus, we can predict that in the design of photocatalytic heterojunction, Ni-RP is a possible candidate to serve as an efficient HER photocatalyst in wide pH environments.

3.3 Water adsorption comparison

Besides thermodynamic driving force, many other factors such as microstructures at the micro or nanoscale, adsorption energy, surface/interface morphological properties, and coupling effects with cocatalyst components also have significant influence on photocatalytic performance. (Li et al., 2015). For a better explanation of the reaction mechanism and the prediction of the catalytic activity of a potential photocatalyst, the adsorption of reactants and the reaction energy should also be included. The complex charge-carrier dynamics as well as surface-reaction interactions should be fully investigated since they can largely affect the apparent quantum efficiency in multi-stage heterogeneous photocatalytic reactions. (Li et al., 2016).

The H₂O adsorption energy has a great influence on interfacial charge transfer, and decides the aggregation extent of H2O molecules near the catalytic active sites, which further triggers the concentration effect and results in more H* generated from water splitting. (Wang F. et al., 2020). The adsorption strength of H₂O molecules is greatly affected by the structural stability of single atoms anchored catalytic surface. In particular, the interactions between the anchored SA-TMs and supporting P atoms have been investigated regarding the iso-surface of charge density difference (Supplementary Figure S6) and the Mulliken charge (Supplementary Table S1). The RP stabilized SA-atoms have an obvious electron exchange with coordinated P atoms. Notably, we discover that the Fe atoms (0.27 e) have a stronger interaction with surrounding P atoms than those of Co-RP (0.24 e) and Ni-RP (0.18 e) counterparts. The stable anchoring site on the RP surface of SA-Cu is different from the other three TMs-RP, which is not in the center of the hexatomic ring formed by P atoms. SA-Cu shows a relatively larger average net charge loss of 0.30 e. The electron flow

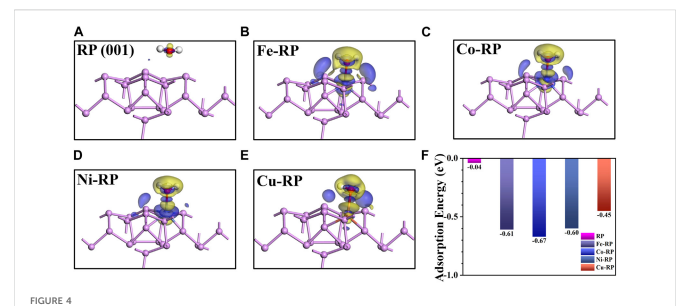
between the stabilized SA-TM atoms and supporting atoms indicates the modification of electronic structures on the RP surface, which verifies the feasibility of doping TMs for photocatalytic activity modulation.

From the charge density difference diagram in Figures 4A-E, H₂O molecules have a much stronger interaction with TMs-RP when compared to pristine RP surfaces. We can see the electron loss in active metals SA sites and adsorbed O atoms (from adsorbed H₂O). Then, the electrons aggregate between TM and H₂O with the formation of an obvious adsorption bond. According to the calculated H₂O adsorption energy values in Figure 4F, H₂O adsorption processes in pristine-RP and TMs-RP are all exothermic, indicating spontaneous adsorption trends. By the loading of TMs SA atoms on the RP surface, the water adsorption capacity has been improved remarkably. Among the four TM-based SACs, Fe-RP (-0.61 eV), Co-RP (-0.67 eV), and Ni-RP (-0.60 eV) exhibit better H₂O affinity than Cu-RP (-0.45 eV). If the H₂O adsorption is too strong, the over-adsorption induces poison effects on the catalytic surface. (Sravan Kumar et al., 2020). Overall, the adsorption energies of Fe-, Co-, Ni-, and Cu-RP are not too negative, indicating less possibility for catalyst poisoning effect during the photocatalysis.

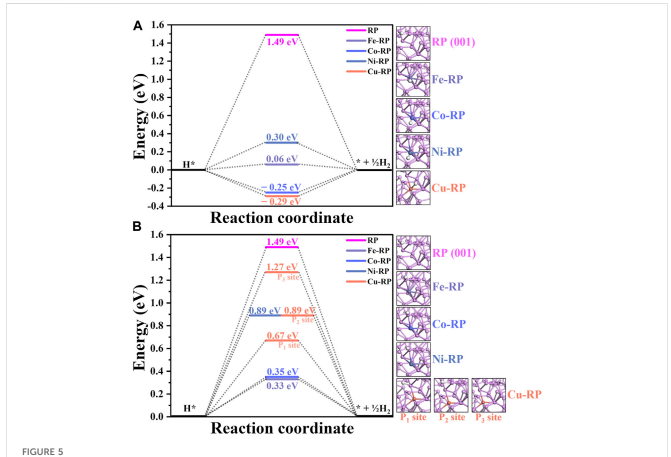
3.4 Reaction energy change of H₂ generation

Besides the water adsorption comparisons, H adsorption is another determinant factor in HER, where both too negative and too positive values are not beneficial for the HER. Based on the adsorption sites, the calculations for H-adsorption reaction energy change are classified into two different types (Figure 5). One considers that H atoms are adsorbed on the SA-metal atoms (Figure 5A), and the other one demonstrates the adsorption of H atoms on the coordinated P atoms to the SA-metal atom (Figure 5B). To guarantee satisfactory computational accuracy, consistent calculation parameters have been set for all the models. For the situations that H atoms adsorbed on SA-metal atoms (H-TMs), the 4 TM SA (Fe-, Co-, Ni-, and Cu-) anchored RP catalysts all have much smaller energy barriers (0.06 eV, -0.25 eV, 0.30 eV, and -0.29 eV, respectively) than that of pristine RP (1.49 eV). As for H adsorbed on coordinated P atoms (H-P-TMs), it can be discovered that the energy barriers of H-P-TMs are slightly larger than those in H-TMs, 0.33 eV (Fe-RP), 0.35 eV (Co-RP), 0.89 eV (Ni-RP), 0.67 eV (Cu-RP, P₁ site), 0.89 eV (Cu-RP, P₂ site), and 1.27 eV (Cu-RP, P_3 site) respectively, but their calculated $\Delta E_{\rm H}$ values are still much less than that of pristine-RP. To sum up, under an acidic environment, the H* is mainly from H* in the solution, where the adsorption energy of H* on the catalyst surface is the main influence factor for HER. Based on the calculated $\Delta E_{\rm H}$, in both situations, H adsorbed on TM atoms (Fe-RP < Co-RP < Cu-RP < Ni-RP < RP) or on coordinated P atoms (Fe-RP < Co-RP < H-P₁- $Cu < Ni-RP = H-P_2-Cu < H-P_3-Cu < RP$), SA TMs-RP have much lower energy barriers than pristine-RP for HER. It is worth noting that in the acidic condition, Fe-RP exhibits the lowest energy HER barrier in both H-adsorption sites discussed above.

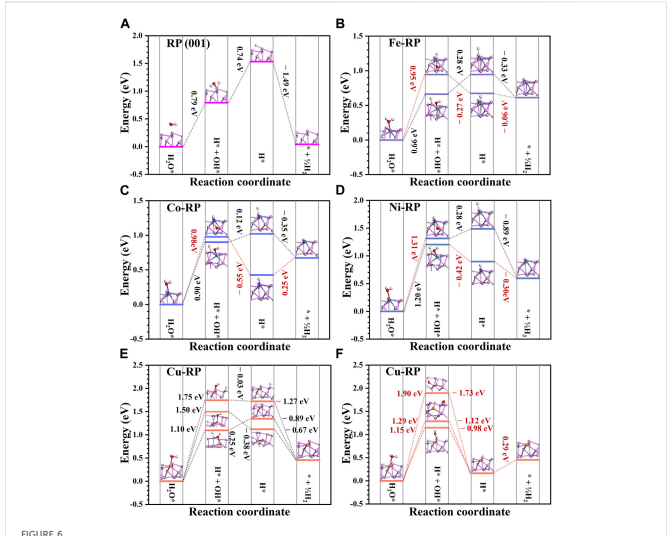
For the photocatalytic mechanism in neutral and alkaline environments, H^* is mainly from water dissociation. The energy



The charge density difference for the net flow of charge between active P/TM atom and O atom in H₂O for **(A)** RP (001), **(B)** Fe-RP, **(C)** Co-RP, **(D)** Ni-RP and **(E)** Cu-RP. The blue color and red color mean the gain and loss of electrons, respectively. **(F)** The comparison of water adsorption energy.



The activation barrier for H adsorption on RP (001) and single atom TM anchored RP surface: (A) H adsorbed on transition metal atoms. (B) H adsorbed on coordinated P atoms.



The reaction energy change of hydrogen generation on (A) RP (001), (B) Fe-RP, (C) Co-RP, (D) Ni-RP, and (E-F) Cu-RP surface. The black dashed lines labeled reaction paths indicate the OH adsorbed on TM atoms and H adsorbed on neighboring P atoms while the red dashed lines labeled paths demonstrate OH adsorbed on coordinated P atoms and H adsorbed on TM atoms during the water dissociation process.

barrier for the dissociation of H2O molecules and the desorption of OH- are important influence factors in reaction energy changes. As shown in Figures 6A-F, the energy change for the whole reaction towards H2 generation is compared. For all the catalysts, the water dissociation step requires the largest energy costs as the ratedetermining step. We have considered two different situations for water dissociation to compare the reaction energy barriers. For the adsorption of OH and H on TM atoms and neighboring P atoms, respectively, the Co-RP and Ni-RP show an energy barrier of 0.90 eV and 1.20 eV, respectively. Different from other TMs-RP types, the anchored single-atom Cu is not symmetrically located in the hexatomic ring of supporting P atoms, which leads to three possible P sites for water dissociations with a water dissociation energy of 1.50 eV, 1.10 eV, and 1.75 eV, respectively (Figure 6E). For water dissociation, Co-RP, Ni-RP, and Cu-RP exhibit higher energy barriers than that of pristine RP (0.79 eV). Notably, Fe-RP shows the smallest energy barrier (0.66 eV) during water dissociation process among all the candidate catalysts. After H₂O dissociation, if the OH is adsorbed on coordinated P atoms and H is adsorbed on TM atoms, the activation energy barriers needed are relatively higher than the former path discussed above, indicating a lower HER efficiency. While the following desorption of OH⁻ becomes much easier on the TMs-RP surface, which requires much lower energy barriers (<0.28 eV) or even becomes spontaneous. In comparison, the pristine RP exhibits a much higher energy cost for the OH⁻ desorption, which leads to the poisoning effect of active sites and lowers the overall photocatalysis performances. Moreover, only Cu-RP and Co-RP (H adsorbed on Co atoms) show endothermic trends for the direct H₂ formation step while other TMs-RP all show the exothermic trend.

For TMs-RP, the energy barriers for OH* desorption are relatively small. In neutral and alkaline environments, since the H* is mainly from the dissociation products (H* and OH*) of adsorbed $\rm H_2O^*$ molecules, the water adsorption energy and dissociation energy are the two main influencing factors for HER. The water adsorption energy follows the order of Co-RP < Fe-RP < Ni-RP < Cu-RP < RP, while the water dissociation energy, which is the largest energy barrier, follows the order of Fe-RP < RP < Co-

RP < Cu-RP < Ni-RP. Less $\rm H_2O$ molecules tend to be gathered around the active catalytic sites on pristine RP surface because its adsorption energy is only -0.04 eV. Thus, although pristine-RP has a relatively small energy barrier for water dissociation, its HER activity will be limited due to the inadequate gathering of reactant $\rm H_2O$ molecules on the catalytic surface. The adsorption energies of water molecules on Fe-RP, Co-RP, and Ni-RP are much lower than that of pristine-RP and Cu-RP, leading to more efficient accumulation of $\rm H_2O$ molecules on these three catalyst surfaces, which will benefit the following water dissociation process. Further taking the order of water dissociation energy into consideration, the activation barriers of Ni-RP and Cu-RP are relatively larger. Overall, in neutral and alkaline conditions, the Fe-RP and Co-RP can potentially offer superior HER performance.

Combining the band alignment and reaction energy, we have proposed photocatalyst candidates for different environments. For the acidic solution, Fe-RP, Co-RP, and Cu-RP are the most promising candidates. For the neutral environments, Fe-RP delivers better performances than other TMs-RP, indicating novel selections to achieve even superior performances than the reported Ni-RP catalysts. (Wang et al., 2022; Jia et al., 2023). In a weak alkaline environment, only Ni-RP is possible to satisfy the requirement of band alignments. As we further consider the electronic structure analysis, the much larger bandgap of RP and Cu-RP significantly lowers the charge separation and transfer process after light excitation, which largely affects their photocatalysis performances. Therefore, we propose that Fe-RP and Co-RP are promising candidates for photocatalysis of H₂ generation.

4 Conclusion

In this work, we have conducted comprehensive investigations on photocatalytic activity regarding the electronic properties, adsorption properties, and reaction energy change for four kinds of SACs by anchoring TM (Fe, Co, Ni, and Cu) single atoms on RP. With the introduction of the single-atom TM anchored on the surface, the sunlight absorption window is extended from red light to a higher NIR region, which potentially improves the utilization rate of solar energy. The electronic structures indicate that Co-RP and Ni-RP show highly catalytic active 3d orbitals to benefit the electron transfer during photocatalysis. Considering the band alignments of photocatalysts, Ni-RP indicates its superiority in supporting HER capability over a wide pH range. Under the acidic environment, Cu-RP, Fe-RP, and Co-RP demonstrate the most preferred proton binding to promote H₂ generation. For the neutral and alkaline solution, all TMs-RP catalysts exhibit much stronger H₂O adsorption than pristine RP, which promotes the following dissociation to supply sufficient proton. In particular, Fe-RP and Co-RP has shown the lowest energy barriers for H₂ generation. Overall, we notice that there is no TMs-RP that has superiority in all pH environments, where different TMs have their specific merits. Based on the comprehensive investigations of different parameters, we think that the TMs-RP possesses great potential and flexibility to achieve efficient H₂ generation in different environments. This work has supplied important theoretical references and opened great opportunities for experimental researchers to further optimize the photocatalytic activity in advanced and novel SACs.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

Author contributions

BH conceived this project and supervised this study. LL and BH carried out the model building and data analysis. MS, TW, QL, BC, CHC, and HHW contributed to the data analysis. LL wrote the first draft of the manuscript. MS, TW, QL, BC, CHC, HHW, and BH contributed to the revision of the article. All the authors approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2023.1197010/full#supplementary-material

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Optimized bisubstrate inhibitors for the actin N-terminal acetyltransferase NAA80

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Acetylation of protein N-termini is one of the most common protein modifications in the eukaryotic cell and is catalyzed by the N-terminal acetyltransferase family of enzymes. The N-terminal acetyltransferase NAA80 is expressed in the animal kingdom and was recently found to specifically N-terminally acetylate actin, which is the main component of the microfilament system. This unique animal cell actin processing is essential for the maintenance of cell integrity and motility. Actin is the only known substrate of NAA80, thus potent inhibitors of NAA80 could prove as important tool compounds to study the crucial roles of actin and how NAA80 regulates this by N-terminal acetylation. Herein we describe a systematic study toward optimizing the peptide part of a bisubstrate-based NAA80 inhibitor comprising of coenzyme A conjugated onto the N-terminus of a tetrapeptide amide via an acetyl linker. By testing various combinations of Asp and Glu which are found at the N-termini of β - and γ -actin, respectively, CoA-Ac-EDDI-NH2 was identified as the best inhibitor with an IC50 value of 120 nM.

KEYWORDS

bisubstrate inhibitor, N-terminal acetylation, acetyltransferase, NAA80, actin, cytoskeleton, coenzyme A

1 Introduction

N-terminal (Nt) acetylation is a highly abundant protein modification, occurring on approximately 80% of the human proteome (Arnesen et al., 2009; Aksnes et al., 2015). The process is catalyzed by Nt acetyltransferases (NATs), transferring an acetyl group from Acetyl CoA to the amino group of the first amino acid in the protein sequence. To date seven NATs have been found in human cells, NatA-NatF, and NatH, having distinctive features in terms of subunit composition, subcellular localization and substrate specificity (Aksnes et al., 2019). Some NATs, NatA, NatB and NatC, have broad substrate pools and act cotranslationally. Other NATs have a very specific substrate pool, such as NatD cotranslationally acetylating histones H2A and H4 (Songkyu et al., 2003; Hole et al., 2011), NatF post-translationally acetylating transmembrane proteins (Van Damme et al., 2011; Aksnes et al., 2015), or NatH/NAA80 post-translationally acetylating actins (Drazic et al., 2018).

The first catalytic mechanism of a human NAT (Evjenth et al., 2012), as well as the first structures of a NAT and NAT-complex bound to its peptide substrate, have been presented (Liszczak et al., 2011; Liszczak et al., 2013; Støve et al., 2016). Taking advantage of this information, we and others have developed peptidic NAT inhibitors that are based on

FIGURE 1
Structure of bisubstrate inhibitors. The bisubstrate inhibitors are constructed as conjugates of Coenzyme A coupled via an acetamide linker to a tetrapeptide amide. The amino acids in the peptide sequence were exchanged systematically to identify improved NAA80 inhibition.

covalently linking mimics of the two substrates of the biochemical reaction, i.e., coenzyme A (CoA) and a short peptide carrying an N-terminal bromoacetyl group (Figure 1) giving a thioether-linked acetyl moiety (Foyn et al., 2013; Støve et al., 2016; Goris et al., 2018; Deng et al., 2021). In these studies it has been found that it is sufficient to use the four amino acids from the N-terminus of the protein substrates to obtain potent bisubstrate inhibitors.

Such bisubstrate inhibitors demonstrate specificity and significant inhibitory potential with IC $_{50}$ and K_i in the low $\mu M/nM$ range. For NatD, increasing the length of the acetyl linker to propionyl has proven beneficial for the design of highly potent inhibitors (Deng et al., 2021). Further, for most NATs, access to bisubstrate inhibitors has led to significant advances in the field by enabling the determination of protein crystal structures (Liszczak et al., 2013; Støve et al., 2016; Hong et al., 2017; Goris et al., 2018; Deng et al., 2020; Deng et al., 2023).

Actin is a major component of the cytoskeleton and cytoskeletal dynamics are important for several cellular activities such as cell motility, division, and intracellular trafficking (Pollard and Cooper, 2009). The cellular activity of actin is connected to its dynamic transition between monomeric (G-actin) and filamentous (F-actin) forms. This is regulated in cells by actin-binding proteins as well as by post-translational modifications. It was established 4 decades ago that animal actins undergo a unique Nt-maturation process (Redman and Rubenstein, 1981; Rubenstein and Martin, 1983). Now we know that this involves the following steps (Arnesen and Aksnes, 2023): a) a general Nt-processing step for all eukaryotic proteins of co-translational N-terminal acetylation (Aksnes et al., 2019) (preceded by methionine excision for muscle actins), b) a unique animal actin-specific post-translational cleavage of the Ntacetylated residue by ACTMAP (Haahr et al., 2022) and finally c) an animal actin-specific post-translational Nt-acetylation by NAA80/NatH to generate acidic actin Nt-termini (Drazic et al., 2018; Goris et al., 2018; Wiame et al., 2018). NAA80 binding to profilin, specifically PFN2, primes the interaction with actin monomers and the acetylation of actin's N-terminus which sticks out from the folded actin monomer (Rebowski et al., 2020; Ree et al., 2020). In human cells lacking NAA80, actin is not Nt-acetylated (near 0% Nt-acetylation) while in the presence of NAA80, actin is acetylated to a near 100% stoichiometry (Drazic et al., 2018; Drazic et al., 2022). This suggests that actin Nt-acetylation is essential for optimal actin functionality in humans. Indeed, human NAA80-KO cells where actin is unacetylated display fragmentation of the Golgi apparatus, altered cytoskeletal organization including decreased G-actin/F-actin ratio, increased F-actin, increased cell size, increased filopodia and lamellopodia and increased cell migration (Aksnes et al., 2018; Drazic et al., 2018; Beigl et al., 2020). The physiological impact of actin Ntacetylation is not fully understood due to the lack of NAA80 KO animal models. However, two brothers carrying a homozygous NAA80 variant resulting in a partial reduction in cellular actin Nt-acetylation were recently presented (Muffels et al., 2021). These individuals showed hearing loss, mild muscle weakness and developmental delay.

Bisubstrate inhibitors of NAT enzymes could potentially be used as tools to study the effects of inhibiting Nt-acetylation to shed more light on the roles of Nt-acetylation in the cell. NAA80 inhibitors could specifically be applied to manipulate cytoskeletal dynamics and to increase cell migration. Herein we describe the optimization of a bisubstrate inhibitor for NAA80.

2 Material and methods

2.1 Synthesis of bisubstrate inhibitors

Bisubstrate inhibitors were synthesised manually or on an Initiator + Alstra (Biotage) automated microwave peptide synthesizer using Fmoc-based solid phase peptide synthesis and ChemMatrix Rink Amide resin (0.44 mmol/g loading) on a 0.25 mmol scale. 2-(6-Chloro-1*H*-benzotriazole-1-yl)-1,1,3,3-tetramethylaminium hexafluorophosphate (HCTU) and

TABLE 1 Analytical data for bisubstrate inhibitors.

Inhibitors	Purity (%) ^a	MS obs. [M–2H] ^{2–}	MS calcd. [M–2H] ^{2–}	MS obs. [M–H] ⁻	MS calcd. [M–H] [–]
CoA-Ac-DDEI-NH ₂	98.4	647.3	647.2	1,295.4	1,295.3
CoA-Ac-DEEL-NH ₂	97.9	654.3	654.2	1,309.3	1,309.3
CoA-Ac-DEEI-NH ₂	95.5	654.3	654.2	1,309.4	1,309.3
CoA-Ac-EEEL-NH ₂	94.1	661.3	661.2	1,323.4	1,323.3
CoA-Ac-EDEI-NH ₂	95.2	654.3	654.2	1,309.3	1,309.3
CoA-Ac-EDEL-NH ₂	94.7	654.3	654.2	1,309.4	1,309.3
CoA-Ac-EEDI-NH ₂	95.1	654.3	654.2	1,309.4	1,309.3
CoA-Ac-EEDL-NH ₂	97.0	654.3	654.2	1,309.4	1,309.3
CoA-Ac-EDDI-NH ₂	93.5	647.3	647.2	1,295.3	1,295.3
CoA-Ac-ESEL-NH ₂	96.8	640.4	640.2	1,281.6	1,281.3
CoA-Ac-EDQL-NH ₂	100	653.8	653.7	1,308.3	1,308.3
CoA-Ac-PDEL-NH ₂	97.3	638.3	638.2	1,277.3	1,277.3

^aBased on RP-HPLC analysis with monitoring at 220 nm

TABLE 2 Inhibitory activity.

TABLE 2 Hillibitory activity.						
Inhibitor	IC ₅₀ (μM) ^a					
CoA-DDDI-NH ₂ ^b	0.38 ± 0.10					
CoA-EEEI-NH ₂ ^b	1.16 ± 0.11					
CoA-MDEL-NH ₂ ^b	1.26 ± 0.10					
CoA-DEEI-NH ₂	1.22 ± 0.33					
CoA-EDEI-NH ₂	0.76 ± 0.18					
CoA-EEDI-NH ₂	0.17 ± 0.04					
CoA-DEEL-NH ₂	4.73 ± 1.43					
CoA-EDEL-NH ₂	0.15 ± 0.02					
CoA-EDQL-NH ₂	0.85 ± 0.05					
CoA-ESEL-NH ₂	11.5 ± 3.23					
CoA-PDEL-NH ₂	13.4 ± 2.24					
CoA-EEDL-NH ₂	1.76 ± 0.52					
CoA-EEEL-NH ₂	1.16 ± 0.12					
CoA-DDEI-NH ₂	2.67 ± 0.67					
CoA-EDDI-NH ₂	0.12 ± 0.05					

 $^{^{\}mathrm{a}}$ All measurements were performed in triplicates. IC $_{50}$ values are given with standard deviation.

N,N-diisopropylethylamine (DIPEA) were used to couple each amino acid (3–5 equivalents) in dimethylformamide (DMF). When the automated microwave peptide synthesizer was used, the coupling process took place with microwave heating at 75 C for 5 min, while manual coupling was performed at room temperature for 30 min. The peptidyl-resin was treated with 20% piperidine in DMF for removal of the Fmoc-protecting group at room temperature for 3 + 10 min. After Fmoc-deprotection of the N-terminal amino acid, the peptidyl-resin was

treated with bromoacetic acid (8 eq.) and N_sN' -diisopropylcarbodiimide (DIC) in DMF for 1 h. The resin was then treated with a mixture of trifluoroacetic acid, triisopropylsilane and water (95:2.5:2.5, v/v/v) for 2 h. The suspension was filtered, and the filtrate was concentrated under reduced pressure until ~5 mL of the solution remained. The crude product was precipitated by adding cold diethyl ether and after removal of the ether layer, the precipitate was triturated with fresh diethyl ether twice. The crude bromoacetyl peptide was dried under

^bData taken from Goris et al. (Goris et al., 2018)

vacuum, purified by semi-preparative RP-HPLC and lyophilised. Purified bromoacetyl peptides and coenzyme A trilithium salt (2 eq.) were dissolved in triethylammonium bicarbonate buffer (1 M, pH 8.5) and left at room temperature overnight. Purification by RP-HPLC and lyophilisation gave the desired CoA-Ac peptides as colorless powders.

2.2 Recombinant protein expression and purification of HsNAA80

HsNAA80/nNat6 (NCBI gene ID:24,142) was subcloned as described by Drazic *et al.* (Drazic *et al.*, 2018) into pETM41 vector. HsNAA80 fused to the maltose binding protein (MBP), was expressed in *Escherichia coli* BL21 star cells at 20°C overnight and lysed with sonication in lysis buffer (300 mM NaCl, 50 mM TrisHCl (pH 8.5), 1 mM DTT, 1x EDTA-free protease mixture). The recombinantly expressed MBP-hsNAA80 was further purified as described by Goris *et al.* (Goris *et al.*, 2018)

2.3 In vitro acetyltransferase activity assay

The enzymatic activity of MBP-hNAA80 was measured using a 5,5′-dithiobis-(2-nitrobenzoic acid, DTNB) assay as described previously by Foyn *et al.* (Foyn et al., 2017) and Drazic *et al.* (Drazic et al., 2018) Briefly, the thiol group exposed in the enzymatic product CoA cleaves DTNB and produces 2-nitro-5-thiobenzonate (TNB⁻) which ionize to TNB² in neutral or alkaline pH and is readily quantified by measuring the absorbance at 412 nm.

2.3.1 Nt-acetyltransferase inhibitor assays

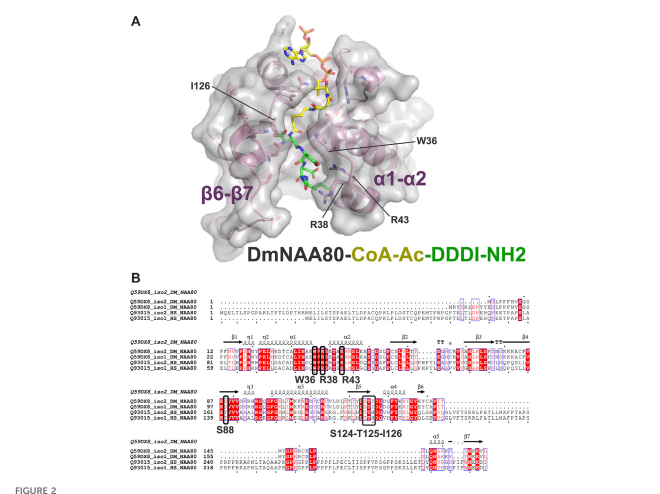
An *in vitro* DTNB assay with enzyme (10–600 nM), 300 μM AcCoA, 300 μM substrate peptide, acetylation buffer (Tris, pH 8.5),

and at least nine different inhibitor concentrations ranging from 0 to 500 μ M was used to calculate the IC₅₀ value for each bisubstrate inhibitor applied in the assay. The reaction was carried out as previously described (Drazic et al., 2018; Goris et al., 2018), and within a timeframe of 15–50 min. All measurements were performed in triplicates.

 IC_{50} values were determined using GraFit 7 software and the results are summarized in Table 2.

3 Results and discussion

The NATs constitute an important class of enzymes and potent inhibitors can prove to be useful as tool compounds to study their roles both in vitro and in vivo. To identify such a tool compound for NAA80, we have in this work optimized the earlier identified bisubstrate inhibitor CoA-Ac-DDDI-NH2 (see Figure 1 for general structure). In our earlier work we tested the in vitro activity of NAA80 toward a broad library of potential substrates such as amino acids, nucleosides, coenzymes, various amines (e.g., serotonin and spermidine), vitamins and a number of 24mer peptides representing the N-terminal part of proteins (Goris et al., 2018). In these substartes, the four N-terminal amino acids were varied while the remaining peptide sequence was kept constant. We found that for all of these potential substrates, NAA80 only acetylated three peptides, MDEL24, DDDI24 and EEEI24. DDDI24 and EEEI24 correspond to the N-terminal part of β- and γ-actin in their cytosolic processed forms, respectively, while MDEL24 represents the unprocessed N-terminal part of the protein p65. The level of product formation for actylation of MDEL24, DDDI24 and EEEI24 was determined to be 119 \pm 5.61 μ M, 50.5 \pm 1.0 μ M, and 42.4 \pm 0.95 µM, respectively (Goris et al., 2018). In our inhibitor design, we decided to continue using a tetrapeptide for the protein-mimicking part as previous studies have shown that these are the most important



Structure of the binary *Drosophila melanogaster* (DM)NAA80/CoA-Ac-DDDI-NH₄ complex and similarities between *Drosophila* and human NAA80 protein sequences. (A) The structure of DmNaa80 is represented with transparent grey surface and pink secondary structures. Side chains involved in binding of the peptide is indicated with sticks, while the bisubstrate inhibitor is highlighted as colored sticks. Key regions ($\alpha 1-\alpha 2$, $\beta 6-\beta 7$) and amino acids (W36, R38, R43, and I126) involved in the peptide binding of the inhibitor are labelled. (B) Sequence alignment of DmNaA80 and *Homo sapiens* (Hs) NAA80. The blue boxes represent sequence conservation while highly conserved residues are shown in red. Strickly conserved residues are white on a red background. Important residues directly involved in peptide inhibitor binding are marked in bold black and with black boxes. Secondary structures and sequence numbering are shown above for the DmNaA80 iso2 sequence. The alignment was made using Clustal T-Coffee and ESPript 3.0.

residues for inhibitory activity (Liszczak et al., 2011; Foyn et al., 2013; Liszczak et al., 2013; Støve et al., 2016; Hong et al., 2017; Goris et al., 2018; Deng et al., 2020; Deng et al., 2021; Deng et al., 2023). Co-crystal structures of NAT-bisubstrate inhibitor complexes typically show that these four amino acids are most important for protein-inhibitor binding interactions. All inhibitors were prepared with the C-terminal carboxylic acid capped as amides to avoid the negative charge of a C-terminal carboxylic acid and better mimic a protein N-terminal.

The peptide part of all inhibitors was prepared using Fmoc-based solid phase peptide synthesis where the C-terminal amino acid was loaded onto a Rink amide resin (Scheme 1). After removal of the Fmocgroup of the N-terminal amino acid, coupling with bromoacetic acid introduced the acetyl-linker. The bromoacetyl peptide was then deprotected and cleaved from the resin and purified by RP-HPLC. Next, triethylammonium bicarbonate buffer (pH 8.5) was used to facilitate the conjugation of CoA to the bromoacetyl peptide. The final bisubstrate inhibitors were purified by RP-HPLC and characterized by MS and NMR to confirm their structures (see Table 1; Supplementary Table S1).

We have already reported that bisubstrate inhibitors based on the N-terminus of processed β - and γ -actin (DDDI and EEEI) display IC_{50} values of 0.38 and 1.16 μM , respectively (Table 2). The difference in inhibitory potency was somewhat higher than was expected based on the small difference in how well NAA80 acetylated the 24-mer peptides DDDI $_{24}$ and EEEI $_{24}$ as their four N-terminal amino acids. Surprisingly, while the MDEL $_{24}$ peptide showed a higher degree of acetylation than the DDDI $_{24}$ and EEEI $_{24}$ peptides when we screened for potential substrates, we found that the inhibitor based on the MDEL sequence was less potent and displayed an IC_{50} value of 1.26 μM .

To elucidate inhibitor binding we have previously solved the crystal structure of DmNAA80 bound to a DDDI bisubstrate inhibitor (Goris et al., 2018). This structure revealed that mainly the $\alpha 1$ - $\alpha 2$ region of NAA80 contributed to peptide binding in the inhibitor, while the $\beta 6$ - $\beta 7$ loop which is important for peptide binding in many of the other NATs, is slightly shifted away from the peptide without contributing to inhibitor binding (Figure 2A). More specifically, D1 of the inhibitor was stabilised by a hydrogen bond from the backbone oxygen of D1 to the backbone nitrogen in S88, while the D1 side chain was stabilized by hydrogen

bonds to the side chain of S124, and through a water-mediated hydrogen bond to T125 and I126. D2 of the inhibitor formed a hydrogen bond with W36 and a salt bridge with R43. Further, D3 made a water-mediated contact with R43 and hydrogen bonds with S46 and S48 and finally, the backbone oxygen of I4 formed a hydrogen bonds with R38. As can be seen from the multiple sequence alignments of DmNAA80 and hNAA80 (Figure 2B), most of the key contributing residues to binding in the DmNAA80-DDDI structure and hNAA80-profilin-actin ternary complex (W36, R43, R38 in DmNAA80 iso2 (Goris et al., 2018) corresponding to W105, R107 and R112 in hNAA80 iso 2 (Rebowski et al., 2020) is conserved. However, a key difference between the two is the extended region around the $\beta6-\beta7$ loop in hNAA80, a region that has shown importance for substrate binding in other NATs (Liszczak et al., 2011; Liszczak et al., 2013; Støve et al., 2016; Hong et al., 2017; Goris et al., 2018), and likely further contributes to optimimal binding of the peptide part of the bisubstrate analogue and increased potency of these inhibitors.

Since the DDDI-inhibitor was found to be significantly more potent that the EEEI-inhibitor, we decided to investigate whether the introduction of Asp into the EEEI sequence would increase inhibitory activity. It turned out that the introduction of Asp in position three was the most beneficial and the inhibitor based on the EEDI sequence showed almost ten-fold higher inhibitor activity compared to EEEI (Table 2). The EEDI sequence also proved to outperform DDDI and was found to inhibit NAA80 with an IC $_{50}$ of 0.17 μ M. Inhibitors based on the DEEI and EDEI sequences gave comparable or slightly improved IC $_{50}$ values respectively compared to the EEEI inhibitor.

We also tested replacing the Ile residue with a Leu and found that the EEEI and EEEL sequences gave inhibitors with equal potency. On the other hand, the DEEL and EEDL sequences gave however a loss of activity compared to DEEI and EEDI while the inhibitor based on EDEL showed a four-to five-fold increase in potency compared to EDEI. Replacing Glu in position three of the EDEL sequence with a Gln residue led to an increase in IC50 from 0.15 μM to 0.85 μM and the replacement of the Asp residue with Ser led to a dramatic loss of potency. Interestingly, a PDEL sequence proved to give only slightly lower inhibitory potency compared to the ESEL sequence.

The introduction of two Asp residues into the EEEI sequence gave inhibitors with a large difference in inhibitory potency as the DDEI sequence gave an IC $_{50}$ of 2.67 μ M, where CoA-Ac-EDDI-NH $_2$ turned out to be the most potent inhibitor with an IC $_{50}$ of 0.12 μ M.

The NAA80/NatH is a particular case among the NAT enzymes in having actin as its sole substrate. The various cytoplasmic and muscle actin substrates all have different combinations of Asp and Glu in the first N-terminal positions (Aksnes et al., 2019). Here, we show that using specific combinations of Asp and Glu in the peptide sequence makes us able to identify a highly potent NAA80 inhibitor. In conclusion, we have found that for NAA80, the choice of acidic residue at different positions of the peptide part of bisubstrate inhibitors is important and varying these has led to an optimized inhibitor with up to 3 fold higher inhibitory activity compared to those previously described. The CoA-Ac-EDDI-NH2 was the most potent NAA80 bisubstrate inhibitor with an IC50 of 0.12 μ M. This inhibitor should be useful in further *in vivo* studies by micro injection in *Danio rerio* and for cell-based studies by micro injection into human cells to elucidate NAA80 function and the effects upon Nt-acetylation of actin.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Materials, further inquiries can be directed to the corresponding authors.

Author contributions

LMM, MB, SIS and HF designed and performed experiments. TA and BEH designed and supervised experiments. LMM, SIS, TA and BEH prepared figures and wrote the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2023.1202501/full#supplementary-material

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Structure, function, and evolution of metallo- β -lactamases from the B3 subgroup—emerging targets to combat antibiotic resistance

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 β -Lactams are the most widely employed antibiotics in clinical settings due to their broad efficacy and low toxicity. However, since their first use in the 1940s, resistance to β -lactams has proliferated to the point where multi-drug resistant organisms are now one of the greatest threats to global human health. Many bacteria use β -lactamases to inactivate this class of antibiotics via hydrolysis. Although nucleophilic serine-β-lactamases have long been clinically important, most broad-spectrum β -lactamases employ one or two metal ions (likely Zn²⁺) in catalysis. To date, potent and clinically useful inhibitors of these metallo-\betalactamases (MBLs) have not been available, exacerbating their negative impact on healthcare. MBLs are categorised into three subgroups: B1, B2, and B3 MBLs, depending on their sequence similarities, active site structures, interactions with metal ions, and substrate preferences. The majority of MBLs associated with the spread of antibiotic resistance belong to the B1 subgroup. Most characterized B3 MBLs have been discovered in environmental bacteria, but they are increasingly identified in clinical samples. B3-type MBLs display greater diversity in their active sites than other MBLs. Furthermore, at least one of the known B3-type MBLs is inhibited by the serine- β -lactamase inhibitor clavulanic acid, an observation that may promote the design of derivatives active against a broader range of MBLs. In this Mini Review, recent advances in structure-function relationships of B3-type MBLs will be discussed, with a view to inspiring inhibitor development to combat the growing spread of β -lactam resistance.

KEYWORDS

antibiotic resistance, β -Lactamase, Metallo- β -Lactamase superfamily, classification, structure, function, evolution, drug development

1 Introduction

The spread of antibiotic resistance is arguably one of the greatest current threats to global health, with multidrug-resistant pathogenic strains increasingly encountered in clinical settings (World Health Organization, 2014). Concerningly, resistance is also being discovered in environmental bacteria that seemingly have not yet been subjected to the same levels of selective pressure as medically relevant pathogenic microbes (though antibiotics are very widely used in farming) (Miraula et al., 2016; Selleck et al., 2020; Wilson et al., 2021).

One of the primary mechanisms of resistance to β -lactams, the most widely employed class of antibiotics (Bush and Bradford, 2016), is via hydrolysis by β -lactamases, which are divided into four Ambler classes: A, B, C, and D (Bush, 2013). Classes A, C and D are serine-β-lactamases (SBLs), which employ a conserved nucleophilic serine residue during hydrolysis of the β-lactam ring. Class B β-lactamases are metallo-β-lactamases (MBLs) that accommodate one or two zinc ions in their active site, which activate a metal ion-bound hydroxide to initiate β -lactam hydrolysis, though at least some MBLs can employ other divalent metal ions (Bahr et al., 2021). Based on their overall sequences, active site residues, mechanisms, substrate preferences, and phylogenetic relationships, MBLs are further subdivided into three subgroups. While evolutionary analyses indicate that subgroups B1 and B2 form a monophyletic group, B3 MBLs appear to have branched independently, suggesting a convergent evolution to acquire the ability to operate on β-lactam substrates (Hall et al., 2004; Bahr et al., 2021). In addition to being distinct from B1 and B2 MBLs, B3 MBLs also display extensive intra-subgroup diversity, notably with active site variations not observed within the B1 and B2 subgroups (Figures 1, 2) (Hall et al., 2004; Pedroso et al., 2020; Bahr et al., 2021).

The MBL-fold, named after its initial discovery in the MBL BcII (Carfi et al., 1995), has a distinctive $\alpha\beta\beta\alpha$ core fold (Figure 1) and is observed in a wide diversity of bimetallic metallohydrolases (Bahr et al., 2021). Together, these form the wider MBL superfamily, with members including phosphatases (e.g., nucleases, phytases), pesticide-degrading esterases, sulfatases (Barbeyron et al., 1995; Hagelueken et al., 2006), lactonases (Fernandez et al., 2011; Miraula et al., 2016), dehalogenases (Wang et al., 2010), oxidases (Muok et al., 2019) and the true β -lactam-degrading members, which form only a small subset within the MBL superfamily (Bebrone, 2007; Bahr et al., 2021). Some MBL fold enzymes also display non-hydrolytic reactions, such as the ethylmalonic encephalopathy 1 protein (ETHE1) that catalyses the iron and oxygen-dependent oxidation of glutathione persulfide (GSSH) to give persulfite and glutathione (Tan et al., 2017; Au et al., 2021; Vašková et al., 2023). Interestingly, there is emerging evidence that many of these diverse activities are observed promiscuously throughout the family (Miraula et al., 2016; Diene et al., 2019; Lee et al., 2019; Perez-Garcia et al., 2021). Notably, a number of recent studies have reported promiscuous enzymatic activities in different members of the MBL superfamily, including human nucleases or glyoxalases with β-lactamase activity (Miraula et al., 2016; Tan et al., 2017; Diene et al., 2019; Lee et al., 2019; Au et al., 2021; Perez-Garcia et al., 2021). Promiscuous enzymes with the MBL-fold could thus represent an as-yet overlooked reservoir from which novel sources of antibiotic resistance may arise under appropriate evolutionary pressures.

Most Class B MBLs are extended-spectrum β-lactamases (ESBLs) and effectively inactivate representatives from each of the three most important groups of β -lactam antibiotics, *i.e.*, penicillins, cephalosporins and carbapenems (Hou et al., 2017; Sidjabat et al., 2018; Bahr et al., 2021). Note, however, that monobactams, such as aztreonam, are not MBL substrates. The strong activity of MBLs against carbapenems, or so-called 'last-resort' antibiotics, is of particular clinical concern (Hou et al., 2017; Sidjabat et al., 2018; Bahr et al., 2021). Recently, infections from Pseudomonas aeruginosa-mediated carbapenem resistance have proliferated and are now a leading cause of death in critically ill and immunocompromised patients (Amsalu et al., 2021; Thaden et al., 2016). The discovery of a highly efficient B3 MBL, Adelaide Imipenemase (AIM-1) in P. aeruginosa, with catalytic efficiencies for the inactivation of a broad range of representatives from each of the three major classes of βlactam antibiotics that is superior to the most effective B1 MBL known to date, NDM-1, demonstrates that the B3 subgroup of MBLs pose a significant but as-of-yet underestimated risk factor to global healthcare (Yong et al., 2012; Selleck et al., 2016; Hou et al., 2017; McGeary et al., 2017). Moreover, while numerous inhibitors for MBLs have been developed (Mohamed et al., 2011; Hussein et al., 2012; McGeary et al., 2014; Arjomandi et al., 2016; McGeary et al., 2017; Krajnc et al., 2019; Palacios et al., 2020; Wachino et al., 2020; Farley et al., 2021) none have yet been used in clinical applications, further exacerbating the threat of MBLs to global health (Mojica et al., 2022). By contrast, potent clinical inhibitors are available for SBLs including the well-established inhibitors clavulanic acid, tazobactam, sulbactam, as well as the more recently developed avibactam and vaborbactam, all of which are used in combination with a β-lactam antibiotic (Hecker et al., 2015; Bush and Bradford, 2016). However, none of the SBL inhibitors are clinically (Bush and Bradford, 2016) effective against MBLs. A major challenge for the development of inhibitors with high specificity for the true MBLs stems from the inherent structural and mechanistic similarity they share with other enzymes from the MBL superfamily [e.g., glyoxalases and some nucleases such as the human enzymes MBLAC1, CPSF73, and HAGH (Pettinati et al., 2016)] as well as enzymes from other families that accommodate bimetallic metal centres that fulfill essential roles in metabolism [e.g. Purple Acid Phosphatases or Phosphodieterases (Mitić et al., 2006; Azevedo et al., 2014)]. The recent discovery of an MBL from the B3 subgroup, CSR-1 from Cronobacter sakazakii, that is inhibited by clavulanic acid may thus present a promising avenue to modify existing anti serine- β lactamase drugs to become broader spectrum therapeutics (Pedroso et al., 2020).

The majority of characterized B3-type MBLs are from environmental bacteria (Hernandez Valladares et al., 2000; Allen et al., 2009; Miraula et al., 2015; Gudeta et al., 2016a; Gudeta et al., 2016b; Pedroso et al., 2017; Rodríguez et al., 2017; Selleck et al., 2020; Wilson et al., 2021) but are increasingly associated with pathogenic organisms as well (*e.g.*, *C. sakazakii*) (Ullah et al., 1998; Yum et al., 2010; Zhou et al.,

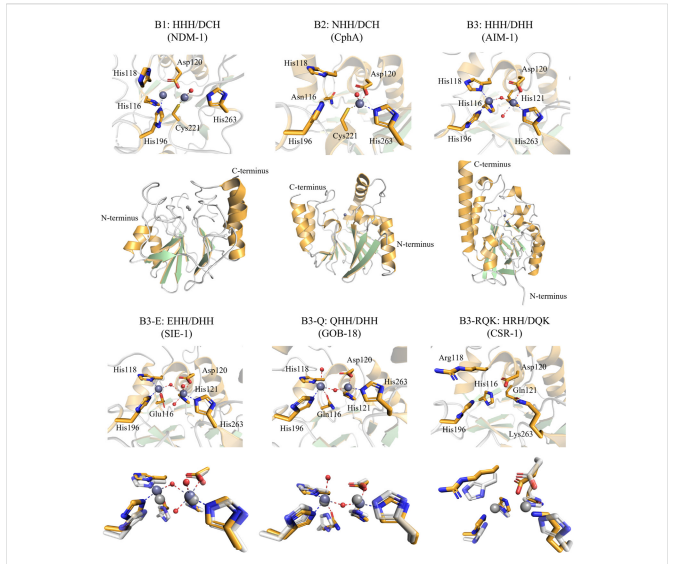


FIGURE 1
Active site and overall structure views of Class B MBLs ("true MBLs") from the B1, B2, and B3 subfamilies. Residues are numbered according to the BBL numbering scheme (Garau et al., 2004). Note that the B1 and B3 MBLs employ two active site metal ions and the B2 MBLs only one metal ion (dark grey spheres: zinc ions). The structures of SIE-1, GOB-18, and CSR-1 were each structurally aligned with AIM-1 (shown in white, 20% transparency, light grey zinc ions) for comparison. The figures were generated using the Pymol visualisation software (Schrödinger, 2015). PDB IDs—NDM-1, a B1 MBL: 3SOZ, CphA, a B2 MBL: 1X8G, and the B3 MBLs: AIM-1: 4P62, SIE-1: 7LUU, GOB-18: 5KOW, CSR-1: 6DN4.

2019). This mini review highlights recent advances in our understanding of this emerging health threat.

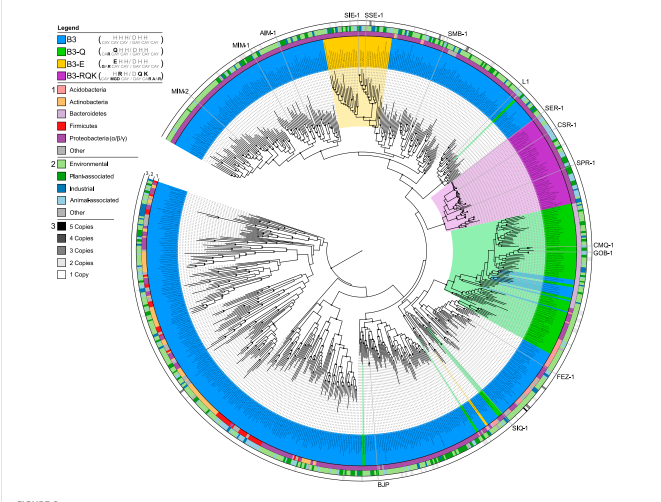
2 Discussion

2.1 Diversity among antibiotic-degrading MBLs

The B1 subgroup has received by far the most attention amongst studies on the Class B MBLs; it contains some of the most notorious agents that confer resistance to antibiotics. B1 subgroup members include enzymes such as NDM-1, IMP-1, and VIM-2 (Crowder et al., 2006), which have been extensively reviewed previously (Mojica et al., 2016; Bahr et al., 2021). B1-type MBLs are

characterised by a conserved HHH/DCH active site motif [i.e., His116, His118, His196 and Asp120, Cys221, His263, as per the BBL numbering scheme (Galleni et al., 2001; Garau et al., 2004)]. The two sets of conserved amino acid side chains provide the ligands that coordinate two zinc ions (i.e., the Zn1 and Zn2 sites, respectively), both of which are required for catalysis (Mitić et al., 2014; Bahr et al., 2021); although it cannot be ruled out that divalent metal ions other than zinc are relevant for MBL activity *in vivo* (Cahill et al., 2016).

The B2-type MBLs constitute the smallest Class B subgroup, of which the best-known member is CphA (Hernandez Valladares et al., 2000). B2 MBLs display the highest degree of substrate selectivity within MBLs, only effectively inactivating carbapenems (Felici et al., 1993; Hernandez Valladares et al., 2000). B2 MBLs are characterised by their NHH/DCH active site motif, corresponding to the Zn1 and Zn2 sites, respectively; however, only one zinc ion, in the Zn2 site, is required for catalysis (Bahr



Phylogenetic analysis of MBLs belonging to the B3 subgroup, highlighting the different active site variants. B3 active site variants are highlighted in different colours according to the figure legend in the top left. The inner circle 1) represents the phylum-level affiliations of bacteria containing B3 MBLs; the middle circle 2) represents the source of the B3-containing bacteria; the outer circle 3) represents B3 MBL gene copy number in each genome. Reproduced with permission from Pedroso et al. (2020).

et al., 2021). Indeed, the presence of a Zn(II) in the Zn1 site leads to inhibition (Mitić et al., 2014; Bottoni et al., 2016; Bahr et al., 2021).

The B3 MBL subgroup contains mostly MBLs detected in environmental organisms and is the most divergent group among the Class B β -lactamases, both from phylogenetic and structural perspectives (Pedroso et al., 2020; Fröhlich et al., 2021).

While the B1 and B2 subgroups seem to have a common evolutionary origin, the B3 subgroup likely evolved independently within the overall MBL superfamily, sharing less than 20% sequence similarity with B1 and B2 MBLs (Hall et al., 2004; Bebrone, 2007; Pedroso et al., 2020; Bahr et al., 2021). Furthermore, the diversity internal to the B3 subgroup seems greater than within the B1 and B2 subgroups, with pairwise similarities being as low as 15% (Pedroso et al., 2020). However, B3-type MBLs possess an active site that can accommodate up two Zn(II) ions, and as for B1 MBLs both Zn(II) ions appear to be needed for full catalytic activity (Selleck et al., 2016). The canonical B3 active site motif is HHH/DHH for

the Zn1 and Zn2 binding sites, respectively. However, unlike the B1 and B2 subgroups there is considerable diversity within the active site motif of the B3 subgroup (Pedroso et al., 2020). Sequence analyses suggest that there are at least four distinct variants that have evolved within the B3 clade, i.e., the canonical form with the HHH/DHH active site motif, and three less abundant forms (Figure 2) (Pedroso et al., 2020). Two of these variants have a substitution of residue 116 of the Zn1 site, replacing the canonical histidine by either a glutamine (Q) or a glutamic acid (E), resulting in the B3-Q (QHH/DHH) or B3-E (EHH/DHH) active site motifs, respectively (Figures 1, 2). The glutamine substitution, in contrast to the glutamate mutation, appears to have occurred independently on multiple occasions within the B3 subfamily (Figure 2) (Pedroso et al., 2020). The least abundant and most intriguing variant, characterised by the B3-RQK active site motif, has three mutations relative to the canonical HHH/DHH motif, i.e., a histidine to arginine of residue 118 of the Zn1 site, and histidine to glutamine and

histidine to lysine in residues 121 and 263 of the Zn2 site, respectively, resulting in the HRH/DQK active site motif (Vella et al., 2013; Pedroso et al., 2020) (Figures 1, 2). This variant appears to form a monophyletic group, having evolved from a single evolutionary event within the B3 MBL subgroup (Figure 2) (Pedroso et al., 2020).

2.2 Structures and catalytic properties of MBLs from the B3 subgroup

a) B3 variants with the canonical HHH/DHH motif. The most studied representatives from the B3 MBL subgroup are L1 from S. maltophilia (Simm et al., 2002; Aitha et al., 2016; Kim et al., 2020) and AIM-1 from P. aeruginosa (Selleck et al., 2016), with the latter posing a major threat to healthcare due to its high catalytic efficiency towards a broad range of β-lactam antibiotics (Supplementary Table S1). The metal ion in the Zn1 site is tetrahedrally coordinated by the three histidine ligands of the active site motif, as well as the oxygen from the hydroxide that in the resting enzyme bridges the two metal ions and which acts as the nucleophile that initiates the hydrolysis of the β -lactam bond. The Zn2 site usually adopts a distorted trigonal bipyramidal geometry where His121 and His263 from the active site motif and the metal ion-bridging nucleophilic hydroxide form the equatorial plane, and Asp120 and an additional water ligand occupy axial positions (Figure 1). Overall, the active site geometry of AIM-1 is similar to that of most of the characterised B1-type MBLs, including that of NDM-1 (King and Strynadka, 2011) or IMP-1 (Concha et al., 2000; Yamaguchi et al., 2021).

MBLs from the B3 subgroup have received far less attention than representatives from the B1 subgroup, largely because they have historically been less often associated with human pathogens. However, they are increasingly being identified in clinical settings and likely pose an underestimated threat to global health (Ullah et al., 1998; Yum et al., 2010; Zhou et al., 2019). For example, AIM-1, originally isolated from a patient from a hospital in Adelaide, Australia, is expressed by bacteria such as *P. aeruginosa* and *K. pneumoniae* (Zhou et al., 2019) and is located on a mobile genetic element (Yong et al., 2012). A recent shotgun genomic sequencing and phylogenetics analysis of wastewater samples from different locations has revealed that, although the gene encoding for AIM-1 may have originated from the non-pathogenic, environmental species *Pseudoxanthomonas mexicana*, it is now widely distributed across the globe (Amsalu et al., 2021).

Furthermore, AIM-1 is highly efficient in inactivating representatives from each of the three major groups of β -lactam antibiotics. In fact, it is considerably more reactive against most of those substrates than NDM-1, one of the most effective agents in antibiotic-resistant isolates around the globe (World Health Organization, 2014; Mojica et al., 2022). In particular, AIM-1 is highly efficient in inactivating "last-resort" carbapenems, with k_{cat}/K_M ratios as high as $18 \text{ s}^{-1}\mu\text{M}^{-1}$, nearly 100-fold higher than the corresponding values reported for NDM-1 (Supplementary Table S1). Mechanistic studies have shown, that like the B1 MBLs, AIM-1 operates most effectively with two Zn(II) bound to the active site (Selleck et al., 2016).

b) B3 variants with the QHH/DHH motif (B3-Q). The best characterised members of the B3-Q MBL variants are the GOB family of MBLs, although several other B3 members have been recovered from environmental soil metagenomes (e.g., LRA-12 or PEDO-1) (Gudeta et al., 2016a; Gudeta et al., 2016b; Rodríguez et al., 2017). GOB-1 was discovered in the opportunistically pathogenic bacterial species *E. meningoseptica* (Horsfall et al., 2011), but numerous GOB variants have since been discovered in *E. meningoseptica* and other *Elizabethkingia* species (Yum et al., 2010; Pedroso et al., 2020; Bahr et al., 2021).

Structural data are available only for one GOB variant, GOB-18 (PDB: 5K0W), and provide evidence that the histidine to glutamine substitution in position116 of the Zn1 motif does not greatly affect the geometry of the active site when compared with that of AIM-1 (Morán-Barrio et al., 2016). However, a noteworthy difference between GOB-18 and AIM-1 is the inversion in the coordination geometries of the Zn1 and Zn2 sites within GOB-18, with Zn2 adopting a tetrahedral and the Zn1 a trigonal bipyramidal geometry (including an axial water ligand) (Figure 1) (Morán-Barrio et al., 2016; Wilson et al., 2021). Similarly to AIM-1, GOB-1 is also effective against a broad range of β-lactam antibiotics, in particular carbapenems, with $k_{\rm cat}/K_{\rm M}$ ratios as high as 8 s⁻¹ μ M⁻¹ (Supplementary Table S1). It is reported that GOB-18 may occur both in mono- and bi-metallic form, similar to the B1-type MBL BsII from Bacillus subtilis, and it may thus be possible that GOB MBLs are active in both states (Morán-Barrio et al., 2007; Lisa et al., 2010). Indeed, the monometallic form may be particularly active against carbapenem substrates, using a mechanism similar to B2-type MBLs that almost exclusively operate on these substrates (Garau et al., 2005; Fonseca et al., 2011).

c) 3 variants with the EHH/DHH motif (B3-E). Only one member of the B3-E variants has been characterised to date, SIE-1 from *Sphingobium indicum* (Wilson et al., 2021), but sequences of other putative B3 MBLs sharing this active site motif have been reported (Pedroso et al., 2020; Bahr et al., 2021). The replacement of the histidine in position 116 of the canonical Zn1 motif by a glutamate does not lead to significant structural changes when compared to AIM-1, preserving a tetrahedral Zn1 and a trigonal bipyramidal Zn2 site (Figure 1) (Wilson et al., 2021). The largest structural difference between the active sites of SIE-1 and canonical B3 MBLs (e.g., AIM-1) or B3-Q (i.e., GOB-18) is that the Zn1-Zn2 distance is longer in SIE-1 (3.75Å) than in AIM-1 (3.48Å) or GOB-18 (3.5Å) (Wilson et al., 2021).

Like other B3 MBLs, SIE-1 is catalytically active against all major classes of β -lactams (Supplementary Table S1). While generally less efficient than the canonical AIM-1 or the B3-Q enzyme GOB-18, SIE-1 has catalytic parameters comparable to those of the B1-type MBL NDM-1 (Wilson et al., 2021). Interestingly, SIE-1 shows increased selectivity toward cephalosporin substrates (with $k_{\rm cat}/K_{\rm M}$ ratios as high as ~1.3 s⁻¹ μ M⁻¹) and is thus more efficient against such substrates than penicillins, an infrequent characteristic amongst B3 MBLs (Wilson et al., 2021) (Supplementary Table S1). From the limited data currently available, it appears that B3-E variants likely employ a mechanism

involving two Zn(II) ions in the active site, similar to AIM-1 (Wilson et al., 2021).

d) B3 variants with the HRH/DQK motif (B3-RQK). The B3-RQK subfamily is the most divergent and least represented among the B3 MBL variants, with substitutions in both the Zn1 and Zn2 sites (Figure 2). To date representatives of the B3-RQK variants have been discovered in and characterised from Serratia proteamaculans (SPR-1) (Vella et al., 2013), C. sakazakii (CSR-1) (Pedroso et al., 2020) and Salmonella enterica (SER-1) (Pedroso et al., 2020). Members of the B3-RQK class are not only unusual because they have the largest number of substitutions in their active sites, but also because two of the canonical histidine residues were replaced by arginine and lysine residues, both very rare ligands in metal ion-dependent enzymes. Due to their high pKa values, these residues are likely to be protonated at pH values optimal for MBL activity, and thus unlikely to coordinate to metal ions effectively. This interpretation is in agreement with initial studies of SPR-1, which indicated that it may have only one Zn(II) bound in the resting state, but that a catalytically active di-Zn(II) centre is formed upon the addition of a substrate (Vella et al., 2013). This substrate-promoted activation mechanism is similar to that proposed for the pesticide-degrading diesterase GpdQ (Hadler et al., 2009; Hadler et al., 2010; Paul et al., 2018; Sharma et al., 2019; Sharma et al., 2020) and indicates an inherent plasticity in the mechanism and potentially the functionality of these enzymes.

Indeed, a high-resolution structure and binding studies of the B3-RQK enzyme CSR-1 reveals that its active site has a strongly decreased metal ion affinity when compared to AIM-1 (Pedroso et al., 2020). Furthermore, while CSR-1 is active against substrates from each of the three major groups of β-lactam antibiotics, it is considerably less efficient than other MBLs (Supplementary Table S1). It has been demonstrated that, at least for some sets of MBLs, that the active site metal ion affinity correlates with catalytic efficiency, in particular for the Zn2 site. In the case of CSR-1 replacement of its arginine in position 118 by the canonical histidine enhances metal ion binding affinity at the Zn1 site with a modest improvement in catalytic efficiency (Supplementary Table S2). However, replacing both the glutamine and lysine residues in the CSR-1 Zn2 site with their canonical histidine counterparts enhances the affinities of both Zn(II) ions and drastically improves the catalytic efficiency to a level comparable to that of other B3 MBLs (Supplementary Table S2). Given that B3-RQK variants evolved from within the B3 MBL subgroup (Figure 2), it is possible that these enzymes indeed represent a functional adaptation, even though no alternative substrates for them have yet been identified.

2.3 Inhibition of B3-RQK MBLs may provide an avenue to combat antibiotic resistance

Given the rise of antibiotic resistance, the lack of clinical inhibitors for MBLs is of pressing concern. A range of SBL inhibitors have been tested to no avail against numerous MBLs, such as penams (e.g., sulbactam and tazobactam),

diazabicyclooctanones (e.g., avibactam and relebactam) (Wang et al., 2016; Lang Pauline et al., 2021), and some boronates (e.g., vaborbactam) (Bush and Bradford, 2016; Cahill et al., 2017; Bahr et al., 2021). Some newer agents such as next-generation boronates (e.g., taniborbactam or xeruborbactam, which are being tested in clinical trials) (Krajnc et al., 2019; Hamrick et al., 2020; Hecker et al., 2020; Liu et al., 2020; Bahr et al., 2021; Brem et al., 2022; Yang et al., 2023), thiol-based compounds (Brem et al., 2016; Tehrani and Martin, 2017) [e.g., captopril (Heinz et al., 2003)], carboxylate-based compounds (Concha et al., 2000; Hinchliffe et al., 2018; Brem et al., 2022), phosphonate-based compounds (Yang et al., 2013; Pemberton et al., 2019), or pyrazoles (Alam, 2022) hold some promise as MBL inhibitors. The recent observation that the commonly used serine-β-lactamase inhibitor clavulanic acid (a clavam) inhibits B3-RQK MBLs may be leveraged to broaden the application scope of this widely used therapeutic. Clavulanic acid is a known inhibitor of serine-β-lactamases (Drawz and Bonomo, 2010), especially those belonging to Classes A and D with K_i values ranging from 20 to 200 μM (Pedroso et al., 2020).

While ineffective against all known MBLs associated with antibiotic resistance in clinical isolates, clavulanic acid inhibits B3-RQK with a K_i of 200–350 μ M (Pedroso et al., 2020). A combination of docking and mutagenesis studies with CSR-1 demonstrated that the lysine residue in position 263 of the Zn2 active site motif may play a crucial role in the binding of clavulanic acid by enabling it to outcompete the catalytically essential Zn(II) in this site (Pedroso et al., 2020). Replacing this lysine with the canonical histidine favours the binding of the metal ion and thus prevents inhibition by clavulanic acid. Considering the overall similarity of the active site geometry in the metal binding pocket of MBLs it may thus be possible to modify clavulanic acid so that its affinity to the Zn2 site will be greater than that of the metal ion.

3 Future perspectives

MBLs have emerged as a major threat to global health, compounded by a lack of clinically suitable inhibitors. The most prevalent MBL subgroup associated with antibiotic resistance is B1, which have highly conserved active site geometries around the catalytically essential Zn(II) ions. However, in recent years an increasing number of MBLs from the B3 subgroup have been identified and characterised. To date, most B3 MBLs are associated with environmental microorganisms that are not pathogenic for humans, but a growing number have been found in more concerning bacteria such as *K. pneumoniae*.

Recent studies have shown that B3 MBLs are evolutionarily related to enzymes with distinct functions, in particular different types of nucleases (Diene et al., 2019), and they have also been identified in several viral genomes. It is thus plausible that B3 MBLs have evolved from a different ancestral functionality and acquired β -lactamase activity in response to environmental pressure. Considering the abundance of such B3 MBLs in diverse microbial organisms it may be only a matter of time before more of these agents accumulate in human pathogens, thus presenting a

new threat to healthcare. It is therefore imperative to think about inhibition strategies to combat them now.

Furthermore, the B3 subgroup is more diverse in terms of active site geometry, mechanism, reactivity, substrate selectivity, and inhibition than the B1 and B2 MBL subgroups. There is a thus a risk that inhibitors developed solely against the currently most important B1 MBL subgroup, which contains the notorious but relatively recently discovered NDM (discovered, 2009), VIM (1999), and IMP (1991) MBLs" to "recently discovered NDM- (discovered, 2009), VIM- (1999), and IMP-type (1991) MBLs, will be become less effective due to a future rise in B3 (and B2) MBLs, against which they may not be active. Interestingly, one active site variant among the B3 MBLs is inhibited by the widely used SBL inhibitor clavulanic acid, by a mechanism involving displacement of the Zn2 metal ion in the active site for which interactions with residue 263 appear to be particularly important. This observation suggests that it may be possible to modify clavulanic acid or related compounds to possess enhanced affinity for the Zn2 site. Along with reports on bicyclic boronates (Cahill et al., 2017; Krajnc et al., 2019), it may be within reach to generate very broad spectrum β-lactamase inhibitors.

Author contributions

SK, SD, PJ, MMP, and AD were responsible for writing the preliminary draft of the manuscript, with support from MTM and GS. CS, LW, and PH were responsible for the critical evaluation of the later versions of the manuscript. GS and MTM were involved in the conceptualization of the manuscript, as well as the final preparation for submission. All authors contributed to the article and approved the submitted version.

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Conflict of interest

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Supplementary material

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Advances in the synthesis and applications of porous carbon materials

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1 Introduction

The progress of human civilization depends on the development of all kinds of materials. The establishment of modern science has led to the rapid development of synthetic materials. However, increasing energy demand and environmental pollution urgently require the search for new materials to solve the energy and environmental crisis.

Carbon is an extremely abundant element in nature and provides the basis for all life on Earth (Li et al., 2008; Toth et al., 2016). The carbon atom has six electrons outside the nucleus, and its outermost electron arrangement is $2s^22p^2$, which shows a strong ability to form covalent bonds (Krueger, 2010). Porous carbon materials have advantages such as chemical stability, low density, high thermal conductivity, high electrical conductivity, and high mechanical strength (Gallo, 2017). Porous carbon materials also have a large specific surface area, adjustable pore size, and functional groups and can be prepared from a wide range of precursors at relatively low cost.

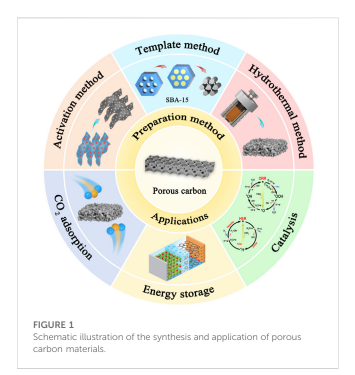
In recent years, a large number of researchers have been devoted to the synthesis and application of porous carbon (Ang, 2019; Liu, 2019; Liu, 2020a; Hwang, 2020; Raj, 2021). Depending on the pore size distribution, the pore structure of carbon materials can be divided into three categories, namely, micropores (pore size <2 nm), mesopores (2 nm < pore size <50 nm), and macropores (pore size >50 nm) (Vu, 2012). The size of the pore structure of porous carbon materials has a significant impact on their performance in practical applications.

Due to these advantages, carbon materials are widely used in the fields of adsorption (He, 2019), catalysis (Dong et al., 2020), and energy storage (Peng, 2019). This paper mainly introduces the synthesis and application of carbon materials and describes the main improvement ideas for current carbon materials (Figure 1). Importantly, the future direction of carbon materials is further discussed.

2 Synthesis methods of carbon materials

2.1 Activation method

Many designs are currently focused on how to increase the specific surface area of carbon materials, which include heat treatment, physical activation, and chemical activation. The physical activation method refers to the use of CO₂, water vapor, O₂, and other gases as activation media for the preparation of activated carbon under high-temperature conditions (Sevilla, 2014; Rashidi, 2019; Ma, 2020). The CO₂ molecules move at a slower rate thermally compared to water vapor activation, resulting in a larger specific surface area and a higher



volume of microporous material. However, the small size and low activation of the activating molecules with a single activator result in a large number of blind pores and an ineffective specific surface area of the carbon material (Kim, 2017).

Chemical activation is the reaction of chemical reagents with the carbon source during pyrolysis, commonly known as KOH, NaOH, H₃PO₄, and ZnCl₂ (Molina, 1995; Lillo, 2003). The main factors influencing the preparation of porous carbon materials by chemical activation include the composition of the precursor material, the activation temperature, the activator, and the impregnation ratio (Maia, 2018). A high impregnation ratio (raw material/activator) allows the formation of activated carbon with a high specific surface area, usually between 1:1 and 5:1 (Rashidi, 2016). KOH is the most commonly used activator, and the pore structure of the prepared material is well developed. Prasankumar converted Tasmanian blue gum tree bark into activated carbons through a simple KOH activationand carbonization method. The generated activated carbons have a hierarchically connected mesoporous structure and a surface area of 971 m² g⁻¹ with an average pore size of 2.2 nm (Thibeorchews, 2022).

Activated carbon is mainly derived from various organic precursors rich in carbon (bitumen, coal, polymers, etc.). Due to the environmentally unfriendly nature of fossil fuels, a large number of activated carbons prepared from biomass as precursors have attracted a great deal of attention in recent years. They are prepared from natural substances such as walnut shell powder (Pang, 2016; Jiang, 2022), banana stem fibers (Taer, 2021), American poplar fruit (Kumar, 2020), bamboo (Khuong, 2022), castor seed (Neme, 2022), and lotus seed shell (Hu, 2018), synthesizing activated carbons with a high specific surface area.

2.2 Template method

Specific surface area and pore size distribution are key factors that affect the properties of carbon materials. The template method

is considered to be an effective method for achieving controlled mesoporous structures. The template method can be divided into the hard template method and the soft-template method. The former involves thorough mixing of the hard template with the precursor materials, carbonization, and subsequent removal of the template. The synthesis of porous carbon materials using various structural silica templates, such as SBA-15 (Yan, 2020) and 3D cubic KIT-6 (Karthikeyan, 2022), has been reported in recent years. The pore size and pore volume of the obtained porous carbon materials can be systematically adjusted by changing the size of the template (Sang, 2011). In addition, porous carbon can also be prepared using CaCO₃, MgO, Mg(OH)₂, magnesium acetate, magnesium citrate, or magnesium gluconate as templates (Wei, 2019; Zhou, 2020), which can be subsequently removed with dilute hydrochloric acid. Liu found a new solvent-free method for the preparation of mesoporous carbon using mesoporous silica KIT-6 as a hard template for the preparation of mesoporous carbon, overcoming the disadvantages of the conventional filling process (Liu, 2019).

The soft-template method involves the self-assembly and cocondensation of a soft template with a precursor to produce a material with a specific structure (Zhang, 2009). The advantages of the soft-template method are that the template does not require subsequent processing and the experimental steps are simple and environmentally friendly. Soft templates include various triblock copolymers, such as F127 and F108. Peng prepared a unique hierarchical porous N-, O-, and S-enriched carbon foam via a combination of a soft-template method, freeze-drying, and chemical etching (Peng, 2019). The structure offers not only an ultra-high specific surface area but also a network of multiple-scale channels.

2.3 Hydrothermal carbonization method

Hydrothermal carbonization is a process in which carbon precursors are gradually hydrolyzed, dehydrated, condensed, and aromatized under high temperature and pressure using water as a solvent and eventually converted into carbon materials. This method is milder and allows for autonomous control of product morphology and better regulation of pore size distribution. In the hydrothermal carbonization process, there are many factors that influence the properties of the carbon material, such as the hydrothermal temperature, the rate of temperature increase, and the holding time. The specific surface area of the carbon material produced is generally low, and the pores are not well developed, so it is often used in combination with activation to obtain porous carbon materials with a high specific surface area.

Liu prepared nitrogen-doped porous carbon materials with a specific surface area of up to 2,864 m 2 g $^{-1}$ and a total pore volume of 1.6 cm 3 g $^{-1}$ by hydrothermal treatment of biomass raw materials and the addition of an activator, KOH, to the aqueous solution, followed by high-temperature pyrolysis and activation (Liu, 2016). Veltri prepared a nitrogen-oxygen co-doped biomass-based carbon material by hydrothermal charring of orange juice with a specific surface area of 1,725 m 2 g $^{-1}$ (Veltri, 2020). The pore structure tended to be reasonable, and the mass fractions of nitrogen and oxygen were as high as 5.65% and 5.38%

3 Applications

3.1 Application of porous carbons in adsorption

The International Panel on Climate Change (IPCC) report shows that the atmospheric concentrations of greenhouse gases (mainly $\rm CO_2$) continue to increase (Lunn, 2021). Porous carbon has a wide range of sources, stable physical and chemical properties, and fast adsorption and desorption rates. It is a kind of $\rm CO_2$ adsorption material with great potential for application.

The carbonized coconut shell was modified with urea at 350° C and activated with $K_2\text{CO}_3$ to produce a nitrogen-doped carbon material with good CO_2 adsorption properties (Yue, 2018). He used polydopamine and melamine as carbon sources and CaCO_3 nanoparticles as templating agents to synthesize porous carbon materials, which showed high CO_2 adsorption capacity and selectivity at room temperature (He, 2019). Pluronic P123 as a soft template was polymerized with D-glucose by the hydrothermal method and activated by CO_2 to produce a porous adsorbent with high microporous content (Nicolae, 2020). The CO_2/N_2 adsorption selectivity of 9 was achieved at 6.00 mmol g^{-1} .

3.2 Application of porous carbon in energy storage

In order to mitigate climate change and environmental pollution caused by excessive use of fossil energy, clean and sustainable alternative energy sources are urgently needed around the world (Weigelt, 2016; Azcárate, 2017). In the past decades, a large number of researchers have been devoted to the development of new types of energy storage (Gondal, 2017; McKone, 2017). Carbon is widely used in energy storage and has the advantages of a large specific surface area, well-developed pores, good electrical conductivity, good electrolyte wetting, high chemical stability, and a wide potential window (Luo. 2020; Kim, 2021). The electrochemical properties of porous carbon electrode materials are a key factor affecting their energy storage properties.

Porous carbon materials are often used as anodes in batteries due to their good electrolyte wetting. The formation of uniform and elastic solid electrolyte interphase (SEI) or the use of tailored electrolytes can improve the stability of the SEI on the carbon surface, thereby increasing the safety and cyclability of the batteries (Fan, 2023; Gu, 2023). The porous carbon used as an electrode in a supercapacitor achieves a high specific capacitance and superior rate capability. Sun prepared nitrogen and sulfur co-doped carbon materials by chemical vapor deposition using magnesium hydroxide as a template, which had a large specific surface area (674 m² g¹¹) and a porous interleaved network with a high level of heteroatom doping (Miao, 2019). As an anode material for Li-ion batteries, it shows a very high reversible capacity and excellent cycling stability.

3.3 Application of porous carbons in catalysis

The development of highly selective, catalytic, stable, green, and economically accessible catalysts is extremely important in industrial production. Carbon materials are often used as

catalysts in ${\rm CO_2}$ electroreduction, oxygen reduction reactions (ORRs), and hydrogen evolution reactions (HERs) due to their abundance of sources and their excellent chemical properties.

Chen proposed an NH3 heat treatment strategy to completely remove pyrrole nitrogen and pyridine nitrogen dopants, and the prepared porous carbon material could efficiently electroreduce CO₂, achieving a CO Faraday efficiency of 95.2% at a current density of -2.84 mA cm⁻² (Dong et al., 2020). Saravanan treated peanut shells using a simple pyrolysis technique assisted by chemical activation and explored the HER properties of peanut shell-derived carbon nanosheets in acidic media (Saravanan, 2019). The nitrogen-doped carbon nanosheets with a high specific surface area and a large number of active sites showed excellent HER catalytic activity in aqueous electrolysis devices (Liu, 2020b). Lai synthesized two-dimensional porous disorderedlayer carbon nanowebs with a large number of N-doped C defects using an aromatic ring as the carbon source and urea as the nitrogen source in a novel molecular design strategy (Lai, 2020). It was shown that carbon edge defects doped with graphitic N atoms could lead to materials exhibiting excellent ORR catalytic properties.

4 Discussions

Due to their high specific surface area, tunable physicochemical properties, low cost, and accessibility, porous carbon materials have shown a wide range of applications in areas such as catalysts, adsorbents, and energy storage. Previously, various methods were used to produce porous carbon materials with different pore structures, but most of the preparation was carried out in the laboratory with complex methods and processes, which made it difficult to meet the requirements of industrial preparation. Moreover, in order to regulate and optimize the structure of porous carbon materials at a more microscopic level and to point the way to the design and preparation of materials, further research on the mechanisms of action of carbon materials in their applications is needed. Thus, with the increasing energy shortage and environmental pollution, the further enhancement of the development of simple, clean, and cost-effective porous carbon materials could make a huge difference in a wider range of areas.

Author contributions

MN conceived and designed the entire review and wrote the manuscript. LZ, YL, and RN edited the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest

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Heterometallic lanthanide complexes with site-specific binding that enable simultaneous visible and NIR-emission

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Macrocyclic lanthanide complexes have become widely developed due to their distinctive luminescence characteristics and wide range of applications in biological imaging. However, systems with sufficient brightness and metal selectivity can be difficult to produce on a molecular scale. Presented herein is the stepwise introduction of differing lanthanide ions in a bis-DO3A/DTPA scaffold to afford three trinuclear bimetallic [Ln₂Ln'] lanthanide complexes with site-specific, controlled binding [(Yb₂Tb), (Eu₂Tb), (Yb₂Eu)]. The complexes display simultaneous emission from all Ln^{III} centers across the visible (Tb^{III}, Eu^{III}) and near infra-red (Yb^{III}) spectrum when excited *via* phenyl ligand sensitization at a wide range of temperatures and are consequently of interest for exploiting imaging in the near infra-red II biological window. Analysis of lifetime data over a range of excitation regimes reveals intermetallic communication between Tb^{III} and Eu^{III} centers and further develops the understanding of multimetallic lanthanide complexes.

KEYWORDS

Lanthanide, luminescence, energy transfer, NIR emission, Macrocycle

1 Introduction

The photophysical properties of lanthanide ions in the common +III oxidation state (Ln^{III}) has become a wide and complex field of research across multiple disciplines and applications, particularly Ln^{III} luminescence for bioimaging (Bünzli and Piguet, 2005; Bünzli, 2010; Monteiro, 2020). Previous works have exploited "windows" of attenuation in the absorption profiles of biological tissues at near-infrared (NIR) wavelengths, allowing deeper penetration of such wavelengths and therefore enhanced imaging (Hemmer et al., 2016; Fan and Zhang, 2019). The Ln^{III} ions are of particular interest due to their characteristic line-like emission across the visible and NIR range, low autofluorescence and photobleaching and high signal-to-noise ratios in comparison to classic organic fluorophores. A caveat to their use is intrinsically poor extinction coefficient as a result of partially-forbidden f-f transitions, which is mitigated by sensitization strategies via organic chromophores (the antenna effect) or d- and f-block metal complexes (Faulkner and Pope, 2003; Natrajan et al., 2009). Furthermore, X-H (X = O, N, C) oscillators are well documented in their ability to

vibrationally quench Ln^{III} excited states due to significant overlap between vibrational overtones of these bonds and emissive Ln^{III} energy levels (Doffek et al., 2012).

This work further develops a previously designed ligand based on a multi-macrocyclic architecture of 1,4,7,10-tetraazacyclododecane-1,4,7-triacetic acid (DO3A) and diethylenetriaminepentaacetic acid (DTPA) (Faulkner and Pope, 2003). Polyamino carboxylate ligands are well known to securely bind lanthanides in a range of environments and have displayed the ability to act as trinuclear heterometallic binding ligands in a site-selective [TbYbTb] arrangement. Herein, we develop the scaffold to include two [LnTbLn] complexes (LnIII = YbIII, EuIII) and a complementary [YbEuYb] species (Figure 1) and further investigate the photophysical capabilities of the ligand across various temperature windows and excitation regimes. The metal selection results in clear Ln^{III}-centred emission across the visible (Tb^{III}, Eu^{III}) and NIR (Yb^{III}) spectrum, which overlaps with both NIR-I (650-950 nm) and NIR-II (1000-1350 nm) imaging windows and therefore invites application in imaging technologies (Foucault-Collet et al., 2013; Jin et al., 2022). The simultaneous response from both LnIII centres has further application as a dual-modal device; the unique magnetic and photophysical behaviour of these metals facilitates use as both MRI contrast agents and optical probes (Rivas et al., 2013; Xu et al., 2013).

Low temperature, solid-state and deuterated media measurements are prioritized to mitigate non-emissive quenching mechanisms. These experiments were also designed to facilitate observation of energy transfer (ET) between metal centers; the relative excited state energies of the chosen metal combinations have been exploited for ET processes in numerous multimetallic Ln^{III} systems (Bispo-Jr et al., 2018; Abad Galán et al., 2021). This work contributes toward the growing library of bimetallic trinuclear Ln^{III} systems that have been studied to elucidate the characteristics of intermetallic communication in discrete molecular complexes (Aboshyan-Sorgho et al., 2012; Zaïm et al., 2014; Tropiano et al., 2015; Maniaki et al., 2023). Finally, the presence of ET in molecular Ln^{III} systems is also useful for imaging applications such as those employing two-photon processes like upconversion (Aboshyan-Sorgho et al., 2011; Nonat et al., 2019).

2 Materials and methods

2.1 Synthesis and characterization

The overall synthesis of target complexes [Yb₂Tb], [Eu₂Tb], and [Yb₂Eu] was adapted and modified from a previously reported literature procedure (Natrajan et al., 2009). Experimental procedures and characterization are detailed on pages S2-S41.

1,4,7,10-tetraazacyclododecane (cyclen) was purchased from CheMatech and used without further purification. All other

reagents and solvents were purchased from Sigma-Aldrich, Fluorochem Ltd. or Apollo Scientific Ltd. and used without further purification. Electrospray +/- (ES-MS) spectra were recorded on a Thermo Orbitrap Exactive Plus mass spectrometer. MALDI-TOF spectra were recorded on a Shimadzu Biotech Axima Confidence mass spectrometer. FT-IR spectra were recorded on a Bruker ALPHA I FT-IR spectrometer. Elemental analysis data were recorder using a Thermo Scientific FlashSmart Elemental Analyzer.

NMR spectra were recorded on a Bruker AVIII HD 500 MHz spectrometer (BBFO inverse probe) in deuterated chloroform, deuterium oxide or deuterated methanol and analyzed using MestReNova 14.1.0. Chemical shifts in parts per million (ppm-δ) are reported relative to residual proton resonances and an internal tetramethylsilane reference. Splitting abbreviations: s: singlet, br. s: broad singlet, d: doublet, dd: doublet of doublets, t: triplet, dt: doublet of triplets, m: multiplet. Blank sections of spectra or those containing solvent resonances are omitted in certain spectra for clarity. Due to complex isomerism between the square and twisted square antiprismatic (SAP ↔ TSAP) forms of multimacrocyclic cyclen compounds, ¹H NMR assignments were often achieved via correlation with 2D COSY, HSQC and HMBC data where possible (Miller et al., 2010; Tircso et al., 2011). ¹H NMR data for compounds containing paramagnetic atoms (YbIII, TbIII, EuIII) were processed using a line broadening/apodization factor of 1.5-5 Hz and baseline corrected using a multipoint baseline correction with a Whittaker, cubic spline or segment algorithm. The chemical shift values of the ¹H resonances in all the lanthanide (III) complexes are reported without assignment due to the complex of the paramagnetic NMR assignment polyaminocarboxylate lanthanide compounds (Sørensen et al., 2017). ¹³C data could not be collected for the same paramagnetic compounds.

Energy minimization of crystal structures was carried out using Avogadro 1.2.0. Compound structures were downloaded as mol2 files from the Cambridge Crystallographic Data Centre (CCDC) and Ln···C distances measured directly from crystallographic data without any further modification to the structure. The superimposed Yb^{III}—Gd^{III} structure was achieved *via* manual manipulation of each structure to minimize steric clash and bonding of the two structures. Bond angles and lengths were maintained during any new bond formation. The auto optimization tool was then used to minimize the energy of the new structure (UFF force field, steepest descent algorithm). Intermetallic measurements were then conducted on this new, minimzed structure.

2.2 Luminescence spectroscopy

Luminescence spectra were recorded on an Edinburgh Instruments FLS1000 Photoluminescence Spectrometer. Solid-state and low temperature spectra were recorded using a cryostat attachment with the sample deposited on a fused silica slide and were the default method of measurement unless stated otherwise. Solution measurements were recorded using a Hellma quartz glass 3.5 mL cuvette with a 1 cm path length and a sample absorbance of 0.1. UV-VIS spectra were recorded using a Mettler-Toledo UV5Bio spectrometer. Samples were excited using a 450 W Xe lamp with a long-pass filter on the detection arm and emission captured by

PMT-900 (visible) and PMT-1700 (NIR) detectors. Any direct comparison of spectra used identical settings; excitation/emission monochromator slit widths and post-collection processing were identical for all. Lifetime measurements were collected using a microsecond flash lamp operating at 40 Hz (${\rm Tb^{III}}$, ${\rm Eu^{III}}$ data) or 100 Hz (${\rm Yb^{III}}$ data). Plotting, fitting and analysis of data was carried out using Origin 2019b. All data were fitted with exponential decay models starting with the fewest terms (mono-, bi-exponential) until sufficiently good fit residuals were achieved. In particular, 2-component fits were always compared against tri-exponential alternatives and found to better fit the data *via* residual, visual, ${\rm Chi}^2$ and ${\it R}^2$ analysis.

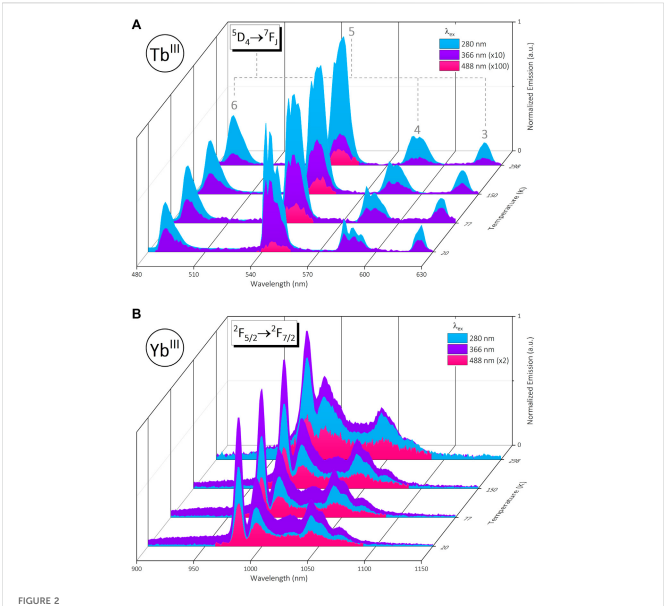
lifetimes from Yb^{III} Shorter-lived emission require consideration of the instrument response function (IRF) of the excitation source, both of which are on a microsecond timescale. An IRF trace was recorded at 100 Hz in line with the procedure detailed by the instrument manufacturer (Edinburgh Instruments) by matching excitation and detection wavelengths ($\lambda = 280 \text{ nm}$) and recording the decay. This was repeated across different temperatures, both with the sample present and separately using milk powder to provide scatter (in both solid and solution-state). The variation in IRF trace between different variables is minimal. Detector response of the (NIR) PMT-1700 was also considered by recording a pseudo-IRF detecting scattered 2λ light from visible excitation ($\lambda_{ex} = 600 \text{ nm}$, $\lambda_{em} = 1200 \text{ nm}$), to provide a similar result. The convolution tool in Origin was used to generate a decay trace with the appropriate IRF taken into consideration, which was then fitted to an exponential decay profile in the same manner as all other lifetime fits. Fitting of Tb^{III} and Eu^{III} signals used the IRF as a benchmark to ensure fitting parameters were only applied after the instrument response had decayed to background and therefore could not contribute to the resulting lifetime.

3 Results

3.1 Photophysical properties of the near infra-red-visible emitting complex [Yb₂Tb]

Initial excitation *via* the phenyl linker ($\lambda_{\rm ex}=280~{\rm nm}$) affords sensitized visible emission from the Tb^{III} metal center $^5{\rm D}_4 \rightarrow ^7{\rm F}_{\rm J}$ (J = 6, 5, 4, 3) transitions. Additional wavelengths selected to directly probe the $^5{\rm D}_3$ ($\lambda_{\rm ex}=366~{\rm nm}$) and $^5{\rm D}_4$ ($\lambda_{\rm ex}=488~{\rm nm}$) energy levels of Tb^{III} also result in emission, however at a lower intensity due to poor extinction coefficients at these wavelengths. Only the primary $^7{\rm F}_5$ transition is visible at $\lambda_{\rm ex}=488~{\rm nm}$. Solid-state variable temperature spectra at 20, 77, 150, and 298 K highlights an increased intensity and resolution of m_j state crystal field splitting at lower temperatures, most notably in the central 545 nm $^5{\rm D}_4 \rightarrow ^7{\rm F}_5$ transition (Figure 2A, Supplementary Figure S2.1).

Simultaneous NIR Yb^{III} emission from the $^2F_{5/2} \rightarrow ^2F_{7/2}$ transition (~980 nm) and the associated 4-fold splitting of the Kramer ground state are present under the same set of excitation regimes across a range of temperatures, with an expected loss in resolution, intensity and broadening of signal at higher temperatures (Figure 2B, Supplementary Figure S2.2). The relationship between emission intensity and excitation wavelength is not concordant between the Yb^{III} and Tb^{III} centers. The signal intensity of the



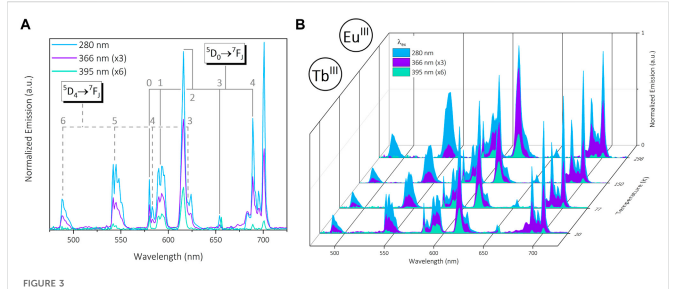
(A): Normalized visible emission of trinuclear bimetallic [Yb₂Tb] under 280 nm (blue), 366 nm (purple) and 488 nm (pink) excitation in the solid state from 20—298 K. (B): Simultaneous normalized NIR emission from [Yb₂Tb] under the same conditions. Data are normalized at each temperature relative to the emission maximum (~545 nm @ λ_{ex} = 280 nm for 2A, ~980 nm @ λ_{ex} = 366 nm for 2B).

Yb^{III} ${}^2F_{7/2}$ transition increases at $\lambda_{ex} = 366$ nm; the intended Tb^{III} 5D_3 state excitation. However, this is not unique to the trinuclear [Yb $_2$ Tb] species as the mono and bimetallic Yb^{III} precursors [Yb] and [Yb] $_2$ DTPA display the same behavior (Supplementary Figures S2.3, S2.4). Inspection of solid-state excitation profiles for the series at 980 nm (Supplementary Figure S2.5) show a broad ligand-centered band with a hypsochromic shift across the series from [Yb] ($\lambda_{max} = 331$ nm) to [Yb $_2$ Tb] ($\lambda_{max} = 311$ nm) and further again for the Tb^{III} center ($\lambda_{em} = 545$ nm, $\lambda_{max} = 296$ nm). Solution-state excitation spectra for the series (Supplementary Figure S2.6) identify this primarily as a solid state effect and present a narrowing of each profile ($\lambda_{max} = 285$ nm) with a similar ~15 nm shift for Tb^{III} excitation ($\lambda_{max} = 270$ nm). UV-VIS absorption data show minimal change in signal when comparing complexes and resemble solution-state profiles (Supplementary Figure S2.7).

Solid-state lifetime measurements of the primary Tb^{III} emission at 545 nm fit a bi-exponential decay profile across all temperatures and $\lambda_{\rm ex}$, determined via analysis of R^2 values and fit residuals (Supplementary Figures S2.8–S2.10). For simplicity, we report the global fluorescence lifetime, τ_n , of each decay component which is an average across variable $\lambda_{\rm ex}$ (280, 366, 488 nm) and temperature (20, 77, 150, and 298 K). This value for the two exponents of Tb^{III} decay are 0.14 \pm 0.01 ms and 0.70 \pm 0.03 ms for τ_1 and τ_2 respectively (fitting after the IRF). The relative contribution of each component varies marginally across the variable excitation and temperature series, but the long-lived τ_2 component remains the most significant (Supplementary Table S2). Calculation of the average lifetime, $\tau_{\rm avg}$ considers the variation in percentage contribution from each τ_n component toward an overall average value (Supplemetary equations S1, S2). Factoring in all data across the excitation and

TABLE 1 Solid-state lifetimes of Ln ^{III} centers at multiple temperatures from the [Yb] complex series including trinuclear bimetallic [Yb ₂ Tb], averaged across variable
excitation wavelengths (298 K: dark red, 150 K: light red, 77 K: dark blue, 20 K: light blue). Tb ^{III} values are calculated average lifetimes from a bi-exponential fit.

Lifetime			Tb ^{III} τ _{avg} (ms)		
Complex		[Yb] (± 0.98 μs)	[Yb] ₂ DTPA (± 1.2 μs)	[Yb ₂ Tb] (± 1.3 μs)	[Yb ₂ Tb] (± 0.03 ms)
Temperature (K)	298	7.6	10	8.1	0.56
	150	7.7	9.9	8.4	0.57
	77	8.0	10	8.7	0.56
	20	8.5	11	9.2	0.51



(A): Normalized visible emission of trinuclear bimetallic $[Eu_2Tb]$ under 280 nm (blue), 366 nm (purple) and 395 nm (green) excitation in the solid state at 20 K. (B): Normalized visible emission from $[Eu_2Tb]$ at 20—298 K across the same set of excitations. Data are normalized at each temperature relative to the emission maximum (~701 nm @ λ_{ex} = 280 nm).

temperature range results in τ_{avg} = 0.55 \pm 0.03 m for the $Tb^{\rm III}$ center (Table 1).

The 980 nm–centered Yb^{III} emission fits a mono-exponential decay following convolution with the IRF signal (Supplemetary Figures S2.11—S2.14) and has a global fluorescence lifetime $\tau = 8.6 \pm 0.4 \, \mu s$ across variable λ_{ex} and temperature. This is comparable to lifetime values for Yb^{III} precursors ($\pm 0.4 \, \mu s$); τ [Yb] = 7.9 μs , τ [Yb]₂DTPA = 10 μs (Table 1).

3.2 Emission behavior of the dual-visible emitting complex [Eu₂Tb]

Insertion of Eu^{III} into the DO3A binding pocket results in strong visible emission (480—700 nm) from the same 7F_J Tb^{III} transitions present in [Yb₂Tb], in addition to the $^5D_0 \rightarrow ^7F_J$ (J = 0–4) transitions from Eu^{III}. As above, excitation of the phenyl linker at $\lambda_{ex}=280$ nm results in simultaneous sensitized emission from the two metal centers (Figure 3A). Direct excitation of the Tb^{III} 5D_3 band ($\lambda_{ex}=366$ nm) again yields emission from both metals proportional to respective excitation profiles. The second direct Tb^{III} excitation (5D_4 ,

 $\lambda_{\rm ex} = 488 \, \rm nm$) employed in [Yb₂Tb] is relatively weak in this complex, meaning no significant emission was observed. Alternatively, direct excitation of the Eu^{III 5}L₆ level ($\lambda_{ex} = 395 \text{ nm}$) can be used to selectively produce EuIII-centered emission and minimize Tb^{III} spectral features, namely, the J = 4, 3 transitions (Figure 3A). Variable temperature emission spectra highlight the consistency of this wavelength-selective visible emission in addition to crystal field splitting at lower temperatures, in particular the Eu^{III 7}J₁ and ⁷J₂ signals (Figure 3B, Supplementary Figure S2.15). The lack of splitting in the $^5D_0 \rightarrow ^7F_0$ transition is indicative of a singular Eu^{III} environment due to the non-degeneracy of both states, as expected from equivalent [EuIII(DO3A)] sites (Binnemans, 2015). Additionally, the hypersensitive ⁷F₂ transition known to be highly dependent on coordination environment remains consistent throughout. Analysis of λ_{ex} against temperature shows an appreciable change in intensity at 701 nm (Eu^{III} $^{7}F_{4}$ state) when changing from sensitized ($\lambda_{ex} = 280 \text{ nm}$) to direct excitation ($\lambda_{ex} = 395 \text{ nm}$) (Supplementary Figure S2.16). However, the relative intensity of the ⁷F₄ signal is strong overall which is in agreement with the square antiprismatic geometry expected of macrocyclic species such as DO3A (Binnemans, 2015).

Life	etime		$Tb^{III} \tau_{avg}$ (ms)		
Complex		[Eu] (± 0.05 ms)	[Eu] ₂ DTPA (± 0.03 ms)	[Eu ₂ Tb] (± 0.02 ms)	[Eu ₂ Tb] (± 0.02 ms)
(K)	298	0.58	0.76	0.15	0.23
ture (J	150	1.1	0.88	0.21	0.35
ıpera!	77	1.2	0.88	0.23	0.34
Ten	20	1.2	0.89	0.24	0.38

TABLE 2 Solid-state lifetimes of Ln^{III} centers at multiple temperatures from the [Eu] complex series, averaged across variable excitation wavelengths (298 K: dark red, 150 K: light red, 77 K: dark blue, 20 K: light blue). Both Eu^{III} and Tb^{III} values are calculated average lifetimes from a bi-exponential fit.

Fitting of the 545 nm Tb^{III} decay results in a bi-exponential fit analogous to the [Yb₂Tb] species ($\tau_1 = 0.11 \pm 0.01$ ms, $\tau_2 = 0.56 \pm 0.07$ ms). The percentage contribution of each lifetime component is more evenly distributed than in the corresponding [Yb₂Tb] complex, resulting in a shorter τ_{avg} value of 0.33 \pm 0.02 ms (Table 2). However, comparison of the longer-lived τ_2 component in both Tb^{III} -containing complexes highlights similarity between the two (Supplementary Table S3).

The Eu^{III} lifetimes were measured by fitting the decay of the primary 615 nm emission ($^5D_0 \rightarrow ^7F_2$) at various temperatures and fit with a bi-exponential equation also (Supplementary Figures S2.17—S2.19) ($\tau_1=82\pm5~\mu s$, $\tau_2=0.36\pm0.04~ms$). The contribution of each component toward τ_{avg} is largely independent of λ_{ex} and differs from Tb^{III} bi-exponential fits as the longer-lived τ_2 component no longer dominates (Supplementary Table S4). Each τ_n component has appreciable significance, yielding a τ_{avg} value of 0.21 \pm 0.02 ms. Comparison of component-weighted τ_{avg} Eu^{III} lifetimes from [Eu] and [Eu]₂DTPA shows they are longer across the series when compared to the final [Eu₂Tb] bimetallic compound ($\tau_{avg}=1.0\pm0.3~ms$ and 0.85 \pm 0.06 ms respectively, Table 2).

Comparison of EuIII precursors [Eu] and [Eu]2DTPA with the trinuclear target species [Eu₂Tb] highlights significant differences in solid-state excitation profiles arising from 615 nm emission (Supplementary Figure S2.20). The DO3A complex [Eu] displays strong absorption bands corresponding to ${}^5D_4 \leftarrow {}^7F_0$ (362 nm), 5D_4 \leftarrow ⁷F₁ (375 nm), ⁵L₆ \leftarrow ⁷F₀ (395 nm), ⁵L₆ \leftarrow ⁷F₁ (400 nm) and ⁵D₂ \leftarrow $^{7}F_{0}$ (465 nm) Eu^{III} transitions. Phenyl ligand absorption (λ_{ex} max ≈280 nm) begins to dominate in the following [Eu]₂DTPA compound, coupled with the loss of the 5D4 band at 375 nm. [Eu₂Tb] follows the trend with increased ligand-centered absorption and further absence of the other 5D4 feature at 362 nm; Eu^{III} series spectra exhibit general solid-state broadening analogous to [Yb2Tb]. Solution-state excitation scans in D2O are ligand-centered across the series, with a marginal shift between samples ($\Delta \lambda_{max} = 20 \text{ nm}$) and a minor 5L_6 feature at 395 nm (Supplementary Figure S2.21) and are in agreement with UV-VIS spectra (Supplementary Figure S2.22).

3.3 Photophysical properties of the near infra-red-visible emitting complex [Yb₂Eu]

Synthesis of the $\{Yb(DO3A)\}_2$ - $\{Eu(DTPA)\}$ complex $[Yb_2Eu]$ results in simultaneous NIR and visible emission from the two metals ($\lambda_{ex} = 280$, 366 and 395 nm) analogous to $[Yb_2Tb]$ and

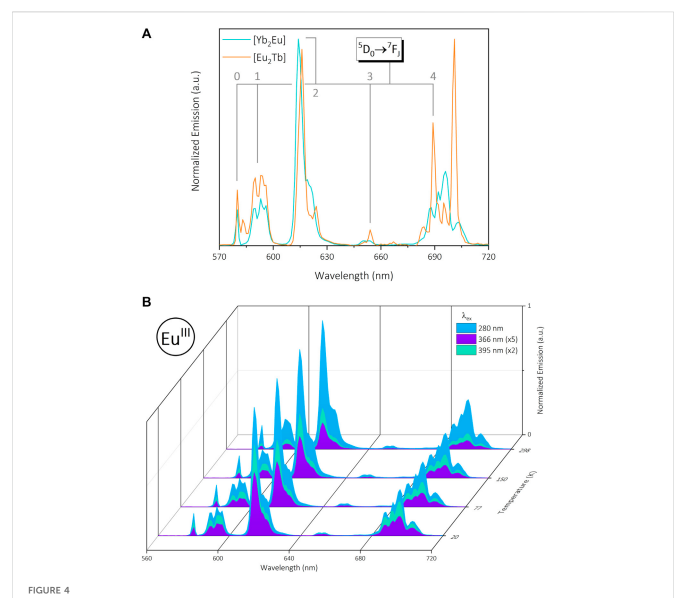
[Eu₂Tb], with an expected decrease in emission intensity at higher temperatures (Supplementary Figures S2.23, S2.24). Excitation at 366 nm was investigated despite the absence of Tb^{III} to generate results comparable with other data sets. The Eu^{III} ⁷F₂ transition displays a minor spectral shift, and the ⁷F₄ presents a change in splitting pattern compared to [Eu₂Tb], both of which are sensitive to coordination environment and suggest EuIII is selectively bound in the DTPA site (Figure 4A). The EuIII center exhibits a greater response at $\lambda_{ex} = 395$ nm compared to [Eu₂Tb], which is consistent across a range of temperatures (Figure 4B). Solid-state excitation spectra (λ_{em} = 615 nm) show a large ligandcentered signal in addition to distinct EuIII bands analogous to [Eu] and rationalize this behavior (Supplementary Figure S2.25). Solution-state excitation spectra of [Yb₂Eu] present ${}^5D_4 \leftarrow {}^7F_0$, ⁷F₁ Eu^{III} excitation signals which are absent in solution measurements of the other Eu^{III} complexes (Supplementary Figure S2.26).

The Yb^{III} emission intensity from the ${}^2F_{5/2} \rightarrow {}^2F_{7/2}$ transition shows an increased temperature-wavelength relationship compared to other Yb^{III} complexes but maintains maximum emission at $\lambda = 366$ nm (Supplementary Figure S2.27). At T = 298 K both ligand and Eu^{III} excitation ($\lambda_{ex} = 280$ nm and 395 nm, respectively) result in similar emission intensity, however the latter begins to dominate at lower temperatures (T ≤ 150 K). Solid and solution-state excitation spectra at 980 nm exhibit the same relationship as [Yb₂Tb] with a \sim 40 nm blue shift between the two metal centers and UV-VIS spectra is comparable to other Eu^{III} species (Supplementary Figure S2.28).

Lifetime data for Eu^{III} emission in [Yb₂Eu] is similar to previous samples with $\tau_1=0.17\pm0.02$ ms, $\tau_2=0.74\pm0.03$ ms across the temperature and excitation range. The individual component contribution reflects the previous DTPA-bound metal (Tb^{III} [Yb₂Tb]), with a clear dominance of the long-lived τ_2 decay (Supplementary Table S5) which gives rise to a longer τ_{avg} value of 0.63 ± 0.09 ms. Yb^{III} lifetimes remain on the same order of magnitude as previous measurements; $\tau=9.8\pm0.6~\mu s$ across variable λ_{ex} and temperature. A summary of temperature-dependent lifetime data for each Ln^{III} across the trinuclear bimetallic series is presented in Table 3 and highlights the general observation of longer lifetimes at lower temperatures.

3.4 Solution-state measurements

Solution-state measurements in water and D_2O result in lifetimes with mono-exponential decay profiles for all metals; the shorter-lived τ_1 component in the bi-exponential $Tb^{\rm III}$ and $Eu^{\rm III}$ solids are not present. Solution measurements are likely



(A): Normalized visible emission of trinuclear bimetallic [Yb₂Eu] (green) and [Eu₂Tb] (orange) under 280 nm excitation in the solid state at 20 K. Data are normalized for each complex relative to the emission maximum. (B): Normalized visible emission from [Yb₂Eu] at 20—298 K under 280 nm (blue), 366 nm (purple) and 395 nm (green) excitation in the solid state (normalized to ~615 nm @ λ_{ex} = 280 nm).

probing an average lifetime arising from minor changes in hydration state and exchange between the square and twisted square antiprismatic (SAP \leftrightarrow TSAP) isomers of the DO3A macrocycles (Miller et al., 2010; Tircso et al., 2011; Nielsen and Sørensen, 2019). Consequently, solution experiments exhibit significantly longer lifetimes on average, with significant gains in deuterated solvent due to reduced energetic overlap of X-D oscillators with emissive Ln^{III} states (Supplementary Figures S2.29—S2.31) (Doffek et al., 2012). The number of bound solvent molecules can be calculated via the inner sphere hydration parameter, q, using a modified Horrocks equation (Supplementary Equation S3, Supplementary Table S1) (Beeby et al., 1999). This calculation takes into account the degree of vibrational quenching by proximate X-H (X = C, N, O) groups for each metal; O-H oscillators contribute significantly for all three, however the significance of N-H groups is relevant

for Eu^{III} only and is negligible in magnitude for Tb^{III} and Yb^{III} systems. Whilst C-H quenching arising from the acetate methylene and DO3A ring groups has a considerable influence on the luminescent lifetimes of near infra-red emitting lanthanides, in the case of Yb^{III} this contribution is small when compared to closely diffusing O-H oscillators and mostly unobserved, due to the long *X*-H ... Ln^{III} distances of these species and a 1/r⁶ dependence for quenching *via* energy transfer. This enables the estimation of *q* by a modified Horrock's equation (Beeby et al., 1999).

Comparison of the luminescence decay in water and D_2O (Table 4) shows q values change depending on both metal choice and binding environment, with the lowest arising from DO3A-bound Yb^{III} in [Yb₂Tb] (q=0.21) and [Yb₂Eu] (q=0.29). The small size of Yb^{III} due to the lanthanide contraction and availability of hard Lewis basic N- and O- donors in the

TABLE 3 Solid-state lifetimes of Ln^{II} centers at multiple temperatures from the $[Ln_2Ln']$ complex series, averaged across variable excitation wavelengths $([Yb_2Tb] = In_1 + In_2 + In_2 + In_3 + In_4 + In$
orange, $[Eu_2Tb] = green, [Yb_2Eu] = blue)$. Both Eu^{III} and Tb^{III} values are calculated average lifetimes from a bi-exponential fit.

Lifetime		Yb ¹¹¹ τ (μs)		$Tb^{III} \tau_{avg}$ (ms)		Eu ^{III} τ _{avg} (ms)	
Complex		[Yb ₂ Tb]	[Yb ₂ Eu]	[Yb ₂ Tb]	[Eu ₂ Tb]	[Eu ₂ Tb]	[Yb ₂ Eu]
(K)	298	8.1	9.1	0.56	0.23	0.15	0.60
rature (F	150	8.4	9.7	0.57	0.35	0.21	0.65
Temperal	77	8.7	10	0.56	0.34	0.23	0.66
Теп	20	9.2	11	0.51	0.38	0.24	0.62
Glo	bal τ	8.6 ± 0.4	9.8 ± 0.6	0.55 ± 0.03	0.33 ± 0.06	0.21 ± 0.02	0.63 ± 0.09

TABLE 4 Solution-state lifetimes of Ln^{III} centers in [Ln₂Ln'] complexes and associated q values for each, at 298 K ($\lambda_{\rm ex}$ = 280 nm, $\lambda_{\rm em}$ = 980 nm (Yb^{III}), 545 nm (Tb^{III}), 615 nm (Eu^{III}).

Complex	[Yb ₂ Tb]		[Eu	Tb]	[Yb ₂ Eu]	
Metal	Yb ^{III}	Tb ^{III}	Eu ^{III}	Tb ^{III}	Yb ^{III}	EuIII
$ au_{H_2O}$ (ms)	0.0019	1.7	0.60	1.0	0.0016	0.58
$ au_{D_2O}$ (ms)	0.0078	3.0	1.9	2.4	0.0080	2.3
q	0.2	1.1	1.0	-	0.3	1.1

octadentate DO3A precludes access to nearby solvent molecules (Faulkner and Pope, 2003). In complex [Yb2Tb], the neighboring DTPA-bound Tb^{III} is more readily accessible due to a larger ionic radius and reduction in kinetic stability of the DTPA binding pocket, resulting in ~1 bound water molecule (Idée et al., 2009; Sørensen and Faulkner, 2018). Similarly, in the complex [Eu₂Tb], q = 1.0 for the DO3A Eu^{III} center as a result of the slightly larger 9 coordinate ionic radius (Sastri et al., 2003) which competes with the steric bulk of the DO3A macrocycle to allow inclusion of 1 solvent donor. In the case of the Tb^{III}(DTPA) centre in [Eu₂Tb], an apparent q value of 2.4 is determined. However, in the case of substantial energy transfer from the assumedly 5D4 excited state of TbIII to the 5D0 state of EuIII occuring, phonon assisted energy transfer processes through the O-H vibrational manifold are in direct competition with those that act to quench the ${}^5\mathrm{D}_4 \to {}^7\mathrm{F}_\mathrm{J}$ transitions. Given that both of these quenching pathways will possess different rate constants, it follows that Horrocks equation is no longer appropriate (Beeby et al., 1999). Analysis of lifetime data for the analogous [Yb₂Eu] complex indicates that any competitive intermolecular energy transfer from the 5D₀ Eu^{III} excited state to the 7F_{5/2} excited state of Yb^{III} is inconsequential, and q of Eu^{III} bound in the DTPA coordination pocket is calculated as 1.1 as expected.

4 Discussion

The potential of this molecular scaffold to facilitate energy transfer between two bound $\mathrm{Ln^{III}}$ ions was a key factor in both the ligand and experimental design. Measurement of the Gd-(DO3A)-aminophenyl acetamide complex [Gd] (analogous to [Yb] and [Eu]) facilitates characterization of the triplet state $\mathrm{T_1}$

of the phenyl linker. The primary Gd ⁶P_{7/2} state is too high in energy to be sensitized, therefore emission of [Gd] is entirely ligandcentered. Solid-state measurements at 298 K display broad fluorescence (λ_{em} = 431 nm), while phosphorescence from the triplet state is observed at 20 K (λ_{em} = 476 nm, Supplementary Figure S2.32). Crucially, this is high enough in energy to sensitize Tb^{III} , Eu^{III} and Yb^{III} emissive states ($T_1 = 21,008 \text{ cm}^{-1}$, Tb^{III} $^5D_4 =$ $20,453 \text{ cm}^{-1}, \text{Eu}^{\text{III}} \, ^5D_0 = 17,227 \text{ cm}^{-1}, \text{Yb}^{\text{III}} \, ^2F_{7/2} = 2924 \text{ cm}^{-1})$ (Carnall et al., 1968b; 1968a; Sastri et al., 2003) (Figure 5). Additionally, the relative energies and stoichiometry of potential donor and acceptor states (2:1 acceptor:donor ratio) of the metal centers are conducive toward intermetallic ET pathways in [Yb₂Tb] (Tb^{III 5}D₄ \rightarrow Yb^{III 2}F_{5/2}) and $[Eu_2Tb]$ $(Tb^{III}\ ^5D_4 \rightarrow Eu^{III}\ ^5D_0)$. The potential $[Yb_2Eu]$ pathway $(Eu^{III} \, ^5D_0 \rightarrow Yb^{III} \, ^2F_{5/2})$ requires too great an energy mismatch to be a competing process, even when considering phonon assistance via the vibrational manifold of O-H oscillators, as is possibly the case in the $Tb^{III} \rightarrow Yb^{III}$ systems.

Existence of energy transfer between bound metals is often strongly correlated with changes in the luminescence lifetime of excited states, where an energetic pathway from the donor excited state to an acceptor results in an overall reduction in lifetime in the former. An observed reduction in lifetime is apparent as temperature increases for each metal site due to increased X-H (X = C, N, O) vibrations and therefore quenching of excited states. Recorded lifetimes for the potential Tb^{III 5}D₄ donor state are largely consistent across a range of excitation bands. There is however a marked decrease in $Tb^{\text{III}} \tau_{avg}$ values between target complexes [Yb₂Tb] and [Eu₂Tb] (0.55 ms vs. 0.33 ms). There is also a significant reduction of Eu^{III} τ_{avg} in $[Eu_2Tb]$ compared to the $\mathrm{Eu^{III}}$ complex series (0.64 ms, ~75%) which is not present in the analogous YbIII(DO3A) acceptor series, suggesting the existence of an relaxation pathway present in [Eu₂Tb] that is absent in [Yb₂Tb]. Furthermore, Table 3 indicates a 48% decrease in Eu^{III} τ_{avg} from [Yb2Eu] to [Eu2Tb] and supports evidence toward a EuIII-TbIII interaction. This is illustrated more clearly when considering the observed rate constants $k_{\rm obs}$ (1/ $\tau_{\rm avg}$), for Eu^{III} emission in various systems. There is an increase in $k_{\rm obs}$ upon addition of Tb $^{\rm III}$ compared to the Eu^{III}-only complex series ([Eu] $k_{\rm obs} = 1000 \ \mu \text{s}^{-1} \ [Eu]_2 \text{DTPA}$ $k_{\rm obs} = 1180 \ \mu \text{s}^{-1} \ [\text{Eu}_2\text{Tb}] \ k_{\rm obs} = 4762 \ \mu \text{s}^{-1}$). Estimation of the potential energy transfer rate constant k_{ET} can be calculated as the difference between the rate radiative decay from EuIII in the presence and absence of a Tb^{III} transfer partner ($k_{ET} = k_{EuTb} - k_{Eu}$), using data from [Eu₂Tb] and [Eu]₂DTPA, respectively. Calculation with τ_{avg} values for these compounds yield a value of $k_{ET} = 3582 \, \mu s^{-1}$. Analogous

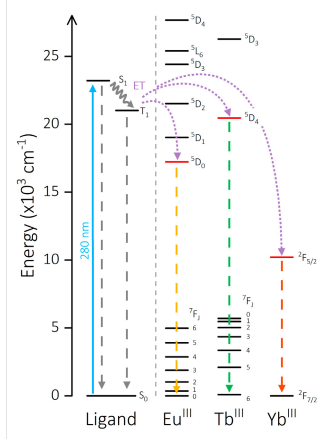
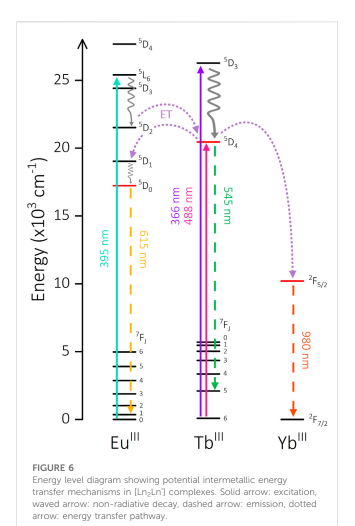


FIGURE 5
Energy level diagram showing the sensitization pathways of [Ln₂Ln'] complexes. Solid arrow: excitation, waved arrow: non-radiative decay, dashed arrow: emission, dotted arrow: energy transfer pathway. Ligand energy levels are calculated from [Gd] measurements. Eu^{III}, Tb^{III}, and Yb^{III} energies are taken from literature for comparison (Carnall et al., 1968a; Carnall et al., 1968b; Sastri et al., 2003).

changes in lifetime are not observed in complexes where either metal is paired with Yb^{III} . Additionally, DO3A-bound Yb^{III} and Eu^{III} complexes show minimal variations in lifetime when paired with a spectroscopically silent Lu^{III} in the central DTPA pocket ([Yb_2Lu] [Eu_2Lu], Supplementary Table S6).

Indeed, the 5D_I energy levels of Tb^{III} (J = 4) and Eu^{III} (J = 2, 1, 0) exhibit appreciable energetic overlap. The 395 nm excitation feature in [Eu₂Tb] populates a Eu^{III 5}L₆ state which is energetically higher than the Tb^{III} 5D_4 level ($\Delta E = 4947 \text{ cm}^{-1}$) and can populate the latter via non-radiative decay to 5D2 and subsequent ET (Figure 6) (Bispo et al., 2018). Additionally, there is the possibility of phonon-assisted ET facilitated by proximate X-H oscillators. The manifold provided by vibrational overtones of O-H and O-D oscillators can provide an alternate energetic pathway to populate ⁷F_I states of Tb^{III} from direct Eu^{III} ⁵L₆ population. Irrespective of these pathways, the emissive states of both Tb^{III} and EuIII are higher in energy than the 7FI states of either metal (Supplementary Figure S2.33) and have been reported to communicate in similar systems (Zaïm et al., 2014). The simultaneous reduction in lifetime of both Eu^{III} and Tb^{III} is exclusive to [Eu2Tb] and is not present in the parent



complexes. The ability of each to act as both donor and acceptor when paired together appears to facilitate a series of ET and possibly back energy transfer (BET) processes between the metal centers.

Despite evidence of solvent interactions and consequent vibrational quenching, solution lifetimes are comparable or often longer than those in the solid state. The additional components in solid-state data likely arise from a combination of SAP ↔ TSAP isomerism and intermolecular quenching between emissive centers in two or more molecules. Solid-state quenching effects appear to play a significant role in the ability of the complex to facilitate energy transfer, as preliminary evidence of Tb^{III}sensitized Yb^{III} emission has previously been reported in solution (Faulkner and Pope, 2003). However, intermetallic distance is another contributing factor and is especially pertinent in solution-state measurements (Sørensen et al., 2017). Single crystal X-ray diffraction structures reported for aminophenyl trifluoromethyl Yb^{III}(DO3A) (CCDC ID: EGOWUV) and bis-aminocarboxyphenyl GdIII(DTPA) (CCDC ID: QEZGIM) complexes with analogous phenyl moieties allow approximate calculation of these distances via superimposition and preliminary energy minimization. Crystal structures were used without modification and manipulated into a feasible geometry

before undergoing optimization in Avogadro to yield an average Yb^{III}–Gd^{III} distance of 11.9 Å (Supplementary Figure S2.34) (Dutta et al., 2006; Pujales-Paradela et al., 2019). Examples of multimetallic architectures that exhibit intermetallic energy transfer often report shorter distances (\leq 10 Å) (Natrajan et al., 2009; Maniaki et al., 2023). The significant r⁻⁶ distance dependence of dipolar ET processes and the innate flexibility of molecular lanthanide systems both act to quench any potential intermetallic communication.

5 Conclusion

A series of three trinuclear bimetallic lanthanide complexes have been synthesized with selective introduction of metals into specific DO3A and DTPA binding sites. All three species exhibit strong sensitized emission when excited via a phenyl linker and represent a broad spectral range, from visible to NIR depending on the metal combination selected. Extensive photophysical measurements investigating the effects of temperature and excitation wavelength revealed communication between EuIII and TbIII centers when closely bound within the same complex, highlighting the potential for energy transfer between metals. The absence of evidence of intermetallic communication in the heterometallics emphasizes the impact of solid-state quenching and solution-state molecular motion on multimetallic lanthanide scaffolds that could facilitate energy transfer. Further work on optimizing the distance between metal centers and varying LnIII selection will be investigated in the future, with the aim of observing energy transfer and therefore accessing more intricate photophysics and applications.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

Author contributions

MET conducted all synthesis, characterization, measurements and analysis for compounds [Yb₂Tb] and [Yb₂Eu]. Initial synthesis

and characterization of [Eu₂Tb] was completed by JH and photophysical measurements finished by MET. LSN, SH, and PP secured funding and provided supervision and discussion. LSN and SF initiated the research and facilitated creation of the project. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2023.1232690/full#supplementary-material

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Compact and modular system architecture for a nano-resonator-mass spectrometer

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Mass measurements in the mega-to giga-Dalton range are essential for the characterization of natural and synthetic nanoparticles, but very challenging to perform using conventional mass spectrometers. Nano-electro-mechanical system (NEMS) based MS has demonstrated unique capabilities for the analysis of ultra-high mass analytes. Yet, system designs to date included constraints transferred from conventional MS instruments, such as ion guides and high vacuum requirements. Encouraged by other reports, we investigated the influence of pressure on the performances of the NEMS sensor and the aerodynamic focusing lens that equipped our first-generation instrument. We thus realized that the NEMS spectrometer could operate at significantly higher pressures than anticipated without compromising particle focusing nor mass measurement quality. Based on these observations, we designed and constructed a new NEMS-MS prototype considerably more compact than our original system, and which features an improved aerodynamic lens alignment concept, yielding superior particle focusing. We evaluated this new prototype by performing nanoparticle deposition to characterize aerodynamic focusing, and mass measurements of calibrated gold nanoparticles samples. The particle capture efficiency showed nearly two orders of magnitude improvement compared to our previous prototype, while operating at two orders of magnitude greater pressure, and without compromising mass resolution.

KEYWORDS

mass spectrometry, NEMS, aerodynamic lens, resonator, single particle

1 Introduction

The characterization of large supra-molecular species is attracting growing interest in biology and analytical chemistry (Keifer et al., 2017; Erdogan et al., 2022). Mass spectrometry (MS) in the MDa to GDa range is especially interesting for the analysis of viral particles and synthetic nanoparticles (Dominguez-Medina et al., 2018; Lai et al., 2021). In this mass range, conventional MS becomes challenged by sample heterogeneity associated with the presence of variant species, chemical modifications, as well as salt or solvent adducts (Rolland and Prell, 2022). This creates highly convoluted m/z patterns that cannot be straightforwardly converted into the original analyte's mass. Several technologies based on single particle mass

determination methods have emerged to circumvent this issue, namely charge detection MS (CDMS) (Jarrold, 2022), nanoelectro-mechanical system-based mass spectrometry (NEMS-MS) (Sage et al., 2015) and mass photometry (Young et al., 2018).

Nano-electro-mechanical system-based mass spectrometry (NEMS-MS) has unique capabilities to analyze ultra-high mass analytes in the MDa to GDa mass range regardless of their charge (Hanay et al., 2012; Sage et al., 2015). Earliest NEMS-based mass spectrometers consisted of modified MS architectures in which the detector was replaced by a nano-resonator (Naik et al., 2009; Hanay et al., 2012). Employing NEMS allowed decreasing the number of components because it works as both detector and analyzer. Moreover, as NEMS-MS does not require charging of the particles, analytes focusing could also rely on particle inertia, while mass is measured directly. These features make NEMS-MS insensitive to mass to charge convolution generated by similar species with differing charge states, enabling a simplified architecture that could ultimately become integrated as tiny instruments. However, as the technique is relatively young (Naik et al., 2009), many features and components still require optimization.

The first NEMS-MS prototype (Naik et al., 2009) consisted of an electrospray ionization source (ESI), a two-stage differentially pumped hexapole ion optics driven at radio frequency, and a NEMS mass sensor localized 2 m below the source. The NEMS resonator was operated at a pressure of 10⁻⁸ mbar and cooled to 40 K. This prototype achieved the first demonstration of NEMSbased MS of single biological molecules. Subsequent research on NEMS-MS focused on simplifying the measurement and the mass spectrometer's architecture, as well as developing more effective methods to focus the analyte on the NEMS mass analyzer/detector. A significant milestone was independently proposed by Hentz and Masselon (2016) and by Malvar et al. (2016). They proposed NEMS-MS systems devoid of ion guides, allowing them to create prototypes with somewhat relaxed pumping requirements. These systems consisted of series of chambers with decreasing pressures: a nebulization stage operating at ambient pressure, a heated capillary inlet with a first pressure drop to 10-100 mbar, a pressure-limiting orifice that could optionally be followed by a series of focusing orifices (aerodynamic lens), and a resonator chamber operating in the 10^{-3} to 10^{-5} mbar regime. These systems allowed decreasing the apparatus size, pumping requirements, and complexity of the system. However, the prototypes were still characterized by modest particle capture efficiency defined as the ratio of the number of detected over emitted particles. In the work by Malvar et al. 1 particle per 5×10^8 was detected, with event rates of ~0.3 particle per minute. Dominguez-Medina et al. (2018) reported the detection of 1 viral particle per 2.6×10^8 with an event rate of 0.8-1.35 particle per minute on a 20 resonators array. Recently, Hannay's group proposed a NEMS-MS system operating under ambient conditions while providing improved particle focusing (Erdogan et al., 2022). As a result, they achieved higher capture efficiency detecting 1 particle per 1.85×10^5 20 nm gold nanoparticles (GNP) and 1 particle per 4.97×10^5 40 nm GNP on one device. However, the atmospheric pressure measurement decreased NEMS performance and mass resolution. Consequently, measurements performed with their prototype on gold nanoparticles and viral particles exhibited substantial mass dispersion.

On the basis of literature reports, we hypothesized that there must exist a favorable operating pressure range that would allow reducing pumping requirements without affecting measurement performance. The goal of the present study was to establish this range through theoretical study and numerical simulation, an eventually demonstrate it experimentally. Ultimately, we applied our findings to develop a new NEMS-MS prototype characterized by decreased pumping requirements, superior focusing performances and improved particle capture efficiency over our previous system (Dominguez-Medina et al., 2018).

2 Materiel and methods

2.1 Working principle of NEMS-MS

Along the development iterations, the core of the NEMS-MS architecture remained unchanged (see Figure 1). It consists of four main parts: a nebulization and desolvation stage, an aerodynamic focusing lens followed by a skimmer, and an array of sensing elements

The NEMS-MS technique analyzes particles from the gas phase, and thus an aerosol must be generated when the analytes are in solution. Two aerosolization methods are typically used: Surface Acoustic Wave Nebulization (SAWN) and nano ElectroSpray Ionization (nESI). The two methods are suitable for biological samples (e.g. viruses, virus-like particles) in small volume (<ml) of buffer solutions. SAWN produces lower kinetic energy droplets, promoting particle sampling through the inlet capillary. Nanoparticles can also be generated using a constant output atomizer, in order to produce larger amount of aerosol required to test system transmission and focusing. The nebulized particle stream passes through a stainless-steel inlet capillary which can be heated up to 250°C in order to dry particles and prevent solvent influence on the mass measurement (Clement et al., 2021). The inlet capillary typically is 250 µm in diameter, and 11 cm long and drives pressure levels in the following chambers of the instrument. Once the solvent excess has been removed, the aerosol must be focused to optimize individual particles capture and detection by the nano-resonator array. An aerodynamic lens is used to perform inertial focusing, producing a narrow particle beam. The longitudinal shape of the particle beam being a cone, the particle number flux (particle/m²/s) decreases with the distance from the lens outlet. Consequently, one of the key parameters of the design is the lens-to-sensor distance, which shall be kept as small as possible. More details about the aerodynamic lens physical principle, simulation and characterization are provided in Section 2.3. Finally, the analyte stream reaches an array of 20 NEMS resonators fabricated from silicon on insulator (SOI) wafers using very large scale (VLSI) integration process (Mile et al., 2010). As the beams are 300 nm wide and their length is about 10 μm , their active area is exceedingly small. An analyte particle reaching a resonator's active surface induces simultaneous shifts in its resonance frequencies, which can be directly related to the mass and position of the landed particle (Sage et al., 2018).

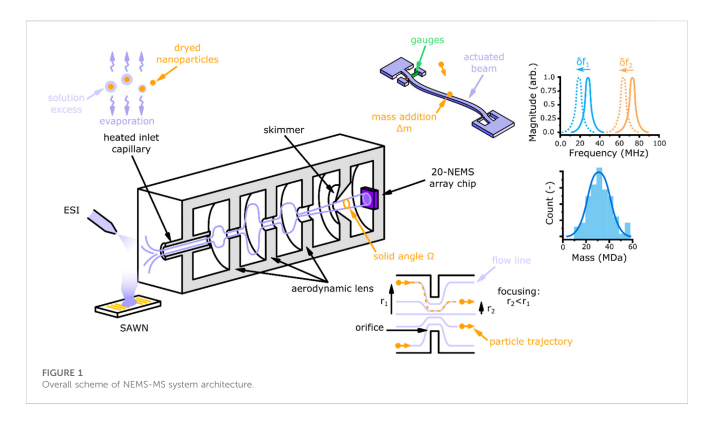


TABLE 1 Length, resonance frequency and quality factors for 3 nano resonators.

NEMS ID	Length [µm]	Mode 1		Mode	2
		f [MHz]	Q [-]	f [MHz]	Q [-]
1	9.04	26.75	4000	73.47	2200
10	8.07	34.80	3600	95.10	1900
20	7.61	44.45	3000	122.00	1700

2.2 NEMS quality factor measurement

2.2.1 Doubly clamped nano resonators

Each NEMS within the array comprises a doubly-clamped vibrating beam electrostatically actuated at its resonance frequency. Two piezoresistive nano gauges near one end of the beam compress and stretch due to the it's displacement, generating a differential signal at the beam's oscillation frequency (see Figure 1). The resonators composing the array were designed with different lengths (see Table 1) in order to be addressed using different resonance frequencies, as shown in Figure 2. They all are 160 nm thick and 300 nm wide and their respective lengths are reported in the Supplementary Material; Section 9.

Resonator-based MS measurements require retrieving the frequency information from each individual resonator. Initial resonance frequencies and phase references are recorded for every resonator in the array (cf. Figure 2). Then, a phase-locked loop (PLL) is locked onto a given resonator to monitor and register frequency data for two modes successively for a given period, called idling time τ_{PLL} , after which it switches to the next resonator. Any

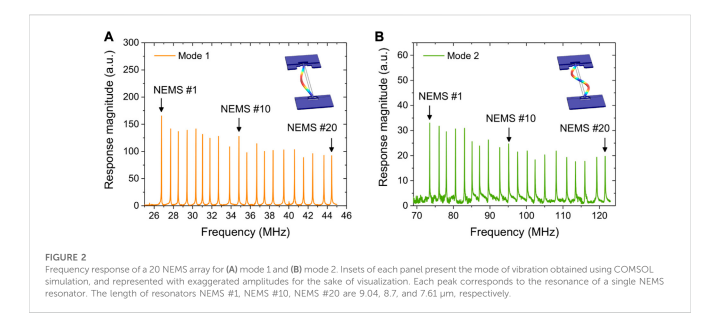
landing particle event results in a mass addition Δm and causes quasi-instantaneous shift in the n-th mode resonance frequency Δf_n . The following relationship describes how Δf_n and Δm are interrelated:

$$\Delta m = M \frac{\Delta f_n}{f_n} \frac{\alpha_n}{\phi_n^2(x)} \tag{1}$$

Where M is the total mass of the beam, f_n the resonance frequency of the n-th mode, x the particle landing position, $\phi_n(x)$ the n-th mode shape and α_n a constant defined by $\alpha_n = -2 \int_{x'=0}^{x'=1} \phi_n^2(x') dx'$. Equation 1 contains two unknowns: the particle mass Δm and its landing location x; thus, a two-equations system must be solved. In practice, the nano-resonator beams are actuated at their first and second modes so the two variables can be derived. The frequency jumps are considered as actual particle landing events when the frequencies shift by amounts larger than 5 times the average frequency noise in each mode (cf. Supplementary Material; Section 5). As a consequence, a crucial parameter for particle mass determination is the frequency stability over the measurement period, which must be as high as possible at the time scale of τ_{PLL} .

2.2.2 Characterizing NEMS for various pressures

The nano-resonators used as sensors can be described by several characteristics, one of which is the quality factor (Q factor), which captures information about energy dissipation in the system. The most significant dissipation for NEMS resonator operating in ambient conditions is caused by viscous damping (Li et al., 2007). To minimize energy losses and therefore maximize the quality factor of the sensor, NEMS are typically operated in a vacuum environment. However, it has been suggested that, as



their critical dimensions approach the mean free path of gas molecules at ambient pressure, the Q factor of nanoscale devices is only marginally reduced while transitioning from high vacuum to atmospheric pressure (Li et al., 2007). Because our goal was to rationalize our system, one possible way entailed relaxing pumping requirements. To evaluate this possibility, we studied the effect of the pressure on NEMS resonance frequencies, their quality factor, and frequency stability. The experimental apparatus used to measure the relevant metrics (i.e., resonance frequency, Q factor and frequency stability) consisted in a vacuum chamber connected to a primary pump and equipped with a 925 micro-Pirani gauge. In order to vary the pressure inside the chamber, a valve was used to create a nitrogen leak allowing to perform measurements in the range $10^{-3} - 10^{3}$ mbar.

2.3 Aerodynamic lens

2.3.1 Inertial focusing

Aerodynamic lenses for aerosol sampling and focusing have been developed since the 1990s (Liu et al., 1995). They use particle drag force and inertia in order to manipulate particles through flow contractions induced by a series of orifices. This produces sudden changes in particle's motion that result in radial particle shifts, for particles having relaxation time higher than the obstacle characteristic time. In aerosol physics, this phenomenon is often characterized using the Stokes number, which is the ratio between particle's inertia and the drag force to which they are subjected, given by:

$$St = \frac{\rho_p d_p^2 C_c(d_p) u}{18\eta L} \tag{2}$$

Where ρ_p is the particle density, d_p its diameter, u the fluid velocity and C_c the Cunningham correction slip coefficient used to correct the drag force expression as a function of the flow regime (molecular, transition, continuum). η is the fluid viscosity and L is

the obstacle characteristic length. When the Stokes number is close to unity, particles deviate from their trajectory due to their inertia but are quickly reattached to a flow line closer to the axis than their previous radial location. This describes how particles should behave when passing through one aerodynamic lens orifice. Since the Stokes number is a function of particle diameter and mass, an aerodynamic lens usually consists of several orifices focusing different particle sizes. Forcing particles to pass through multiple orifices therefore results in a narrower particle beam for a range of particle sizes.

2.3.2 Numerical model

Several authors modelled how nanoparticles behave within an aerodynamic lens (Zhang et al., 2002; Wang et al., 2005a; Abouali et al., 2009). A common approach is based on lagrangian tracking of the particles through the aerodynamic lens, ultimately allowing computation of the particle beam solid angle. COMSOL Multiphysics was used to solve steady, compressible, viscous, laminar Navier-Stokes equations. For computation efficiency's sake, an axisymmetric geometric domain was used. When it comes to boundary conditions, no slip boundary was used for walls, and both the downstream pressure and the throughput mass flow were computed using a macroscopic vacuum system model (cf. Supplementary Material; Section 3). As for the lagrangian part of the simulation, two forces were taken into account: the drag force and the Langevin force used to model brownian motion. The analytical formulations of both forces feature the Cunningham correction slip factor, depending itself on the flow regime (i.e., the Knudsen number) which had to be updated due to the pressure gradient particles are travelling through. The physical model was solved using the velocity Verlet algorithm with an adaptative time step and was written in Python.

2.3.3 Characterization

First, the aerodynamic lens focusing ability was characterized based on polystyrene nanoparticle deposition on a silicon target located at the same position than the detector. The polystyrene 100 nm nanoparticles colloidal solution (Magsphere, Pasadena,

California, United States) concentration was 1×10^{11} particles/ml and was nebulized using nESI at a flowrate of 8 μ l/ min for 30 min. The inlet capillary was heated at 85°C. In order to estimate the diameter of the deposition pattern, the target was observed using binocular loupe and the resulting picture was processed by analyzing pixel intensity profiles (cf. Supplementary Material; Section 7).

The second approach aimed at validating the transmission efficiency of the focusing lens by using the NEMS sensors, and comparing first and second NEMS-MS prototypes characteristics. In this approach, three gold nanoparticles colloidal solutions were investigated. 20 and 40 nm diameter GNP were purchased from BBI Solutions (Crumlin, Wales, UK) and the nm diameter GNP were purchased from Sigma Aldrich (Saint-Louis, Missouri, United States). GNP were sprayed using SAWN or nESI. The mass spectra—Converted into geometric diameter—And event counts per minute and per active NEMS devices were evaluated.

2.4 Pumping requirements

As our overall objective was to decrease the pumping requirements, the design phase included a pressure calculation step. A mass conservation model (cf. Supplementary Material; Section 3) was used to compute the pressure in each chamber based on a given pumping speed. It helped finding design parameters (inlet capillary length and diameter, lens orifice diameters, skimmer diameters) matching NEMS and aerodynamic lens compatible pressure ranges. This model was a keystone in the iterating process for simulating several pumping setups. Importantly, the optimal pressure range for pump performance was also taken into account, as turbomolecular pumps capacity steeply degrades below 10^{-3} mbar.

3 Results and discussion

One of the key parameter to downsize NEMS-MS prototype was the operating pressure: turbo-molecular pumps are large and heavy and require cumbersome vacuum chambers and frame. However, limiting the pressure is necessary to avoid the nano-resonator dampening leading to degradation of the quality factor and thus poorer mass resolution. Furthermore, the operating pressure of the aerodynamic lens, linked to a large extent to the analytical throughput, also plays a role in the quality of the inertial focusing. As a consequence, both of these aspects had to be addressed independently in order to determine the suitable operating pressure range.

3.1 Nano-resonators in moderate vacuum

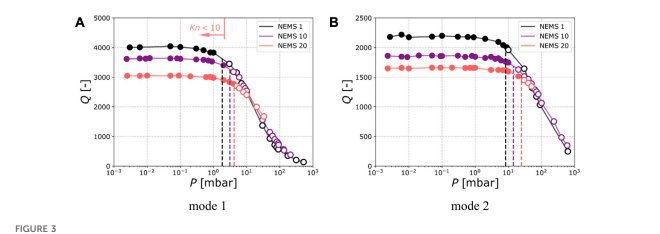
We studied the dependence of the quality factor (Q factor) of several resonators with respect to the operating pressure. In order to cover the various beam dimensions from our 20 resonators arrays, we selected three different beams (i.e. the ones with smallest, largest and median length) and characterized their mechanical parameters—Resonance frequency, quality factor, and frequency stability—As a function of pressure.

Figure 3 displays the Q factors as a function of operating pressure for the three selected resonators. Before reaching high pressures, Q factors are constant and are, for resonators 1, 10 and 20: 4000, 3600 and 3000 for the first mode. For the second mode, their values are 2200, 1900, and 1700, respectively. A loss of 10% of the maximum value (highlighted by dashed lines) was observed at 2, 3, and 4 mbar for the first mode and at 10, 20, and 25 mbar for the second mode for NEMS number 1, 10 and 20 respectively. The trend presented by quality factor as a function of pressure has been previously reported (Li et al., 2007; Gavan et al., 2009), and our results confirm prior observations for our specific devices. A model based on the hydrodynamic function with comparisons to other models was described in (Aoust et al., 2015). The manifestation of the minor effect of the pressure on the quality factor can be given by the flow regime. It can be determined by the Knudsen number $Kn = \lambda/L_c$, which is the ratio between the molecule mean free path λ and a resonator characteristic dimension L_c . In our case, the width of the beams (300 nm) can be considered to estimate the Knudsen regime (Gavan et al., 2009). If the mean free path λ is larger than L_c (when Kn > 10), the probability that the beam encounters a gas molecule is small, leading to a null viscous damping. In this regime, the damping is mainly caused by thermoelastic effects and energy transfer into the support (clamps). In our case, the most significant losses are connected with energy dissipation into the support (cf. Supplementary Material; Section 6). Viscous damping is introduced for a Knudsen number Kn < 10, leading to decrease of the quality factor. The vertical line on the graph Figure 3A was plotted to delimit the relevant Knudsen regimes. For devices with characteristic dimensions in the hundreds of nanometers, this regime shifts toward higher pressure values, and the operating pressure may be increased without sacrificing performance.

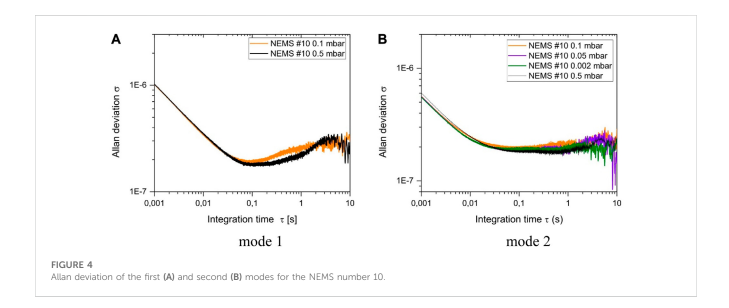
Another parameter that can determine the flow regime is the Weissenberg number $Wi = \tau/T$, which is a ratio of the characteristic time scale T to the relaxation time τ in the medium. It has been shown (Karabacak et al., 2007) that increasing the resonance frequency of nano-resonators allows to reach the molecular flow regime where viscous damping becomes negligible. This may explain why, for the second mode of vibration, the quality factor was less influenced by a comparable increase in pressure than the first mode. Moreover, for the same mode of vibration, smaller resonators (i.e. having a higher frequency) are characterized by a lesser quality factor reduction as a function of pressure, which was also consistent with previous observations (Karabacak et al., 2007).

Frequency stability is a key performance parameter for nanoresonators used in mass spectrometry as the particle masses are deduced from frequency shifts. Lower frequency stability causes higher noise level and degraded mass resolution. The frequency stability was characterized using Allan deviation and was measured as a function of pressure from the closed loop frequency trace of an individual resonator. We show that this parameter remains constant until a pressure of $\sim 0.5\,mbar$ (cf. Figure 4), showing that mass resolution and hence measurement performance remain stable up to this value. Other Allan deviation plots are reported in the Supplementary Material; Section 1.

Our investigations showed that our nano-resonators could operate in a pressure regime as high as 0.5 mbar without sacrificing mass resolution. Moreover, they confirmed that using



Resonance quality factor of the first (A) and second (B) modes for the NEMS number 1, 10 and 20 when operated in at various pressures. This three NEMS have different lengths and thus different resonance frequencies (see Table 1). The dashed vertical lines indicate pressures at which the quality factor loses 10% of its maximum value.



smaller NEMS would allow further reduction of the pumping requirements and thus limit the apparatus footprint.

3.2 Aerodynamic focusing

The ability to operate nano-resonators at a relatively high pressure (or low Knudsen number) is the foundation of the NEMS-MS prototype downsizing. Yet, another important aspect of the instrument depends on pressure: particle transport to the NEMS detector. Therefore, we had to also validate the operation of the aerodynamic lens at higher pressure.

The first NEMS-MS prototype (Dominguez-Medina et al., 2018) integrated an aerodynamic lens which was designed using (Wang et al., 2005b) guidelines. It was optimized to collimate virus-like nanoparticles of diameter in the 100 nm range and unit density. It was experimentally characterized with 45 nm polystyrene particles at an upstream pressure of 10² mbar and a downstream pressure of

 10^{-2} mbar. The resulting deposit at the NEMS location was measured to be 1.5 mm diameter. Since this performance matches the targeted NEMS-MS applications, we decided to retain the same aerodynamic lens design and investigate its focusing abilities at higher pressure.

As described earlier, the selected approach was based on the particle lagrangian tracking through the aerodynamic lens because it could directly provide the particle beam solid angle. Boundary conditions, including mass flow and downstream pressures, were computed using the vacuum system model (cf. Section 2.4; Supplementary Material; Section 3). Ultimately, calculated pressures were compared with actual measurements and found to be in good agreement.

Figures 5A–D present particles trajectories inside the aerodynamic lens computed with an exit pressure of 1.33 mbar for particles of different diameters and a density of 1.06 g/cm³, which corresponds to polystyrene and is close to that of viral samples for which the lens was initially designed. Based on these results, both the

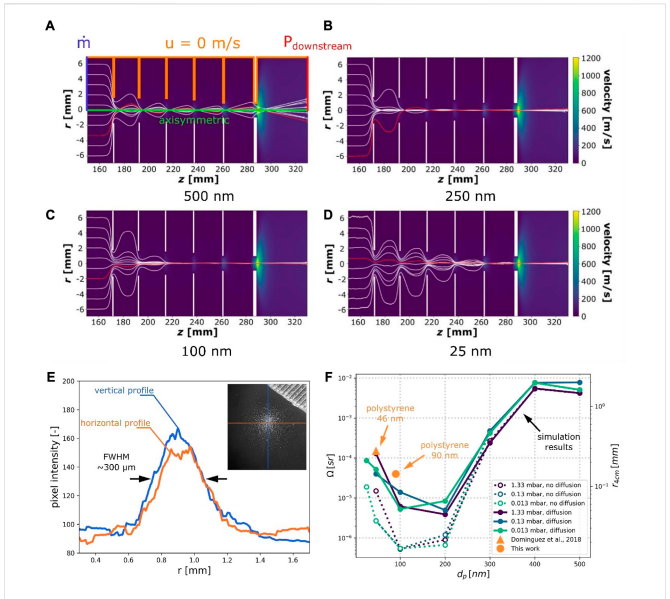
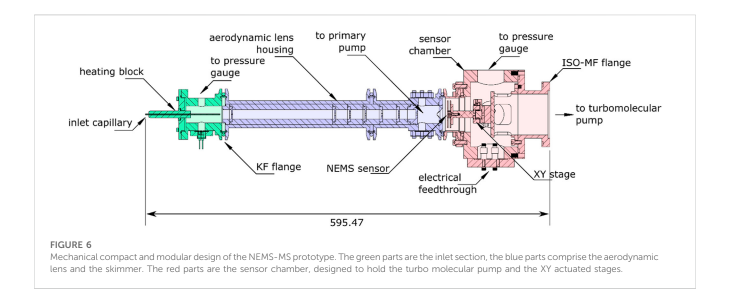


FIGURE 5
Particle trajectories through the lens simulated for different particle diameters: (A) 500 nm, (B) 250 nm, (C) 100 nm and (D) 25 nm. (E) Pixel intensity profiles of the deposit photograph along two axes, located by the orange and blue lines on the inset. (F) Solid angle Ω and particle beam radius at 4 cm downstream the lens (location of the NEMS sensor) computed for several downstream pressures. Orange points represent experimental data and dots are simulation results.

solid angle Ω and the particle beam radius 4 cm downstream the aerodynamic lens (written r_{4cm}) were derived for different particle diameter d_p . The 4 cm distance corresponds to the location of the NEMS array in the new mechanical architecture. The results reported in Figure 5F show how the solid angle behaves depending on particle diameter and that the solid angle remains optimal for the range $100-200\, nm$. Moreover, the addition of the Langevin force used to model diffusion plays a significant role only for particles smaller than $300\, nm$ and yields more realistic predictions when compared to experimental results (orange dots). Finally, within the investigated pressure range $(1.33 \times 10^{-2}-1.33\,$ mbar), the model showed that the impact of diffusion remained relatively constant and pressure had a limited influence on the resulting particle beam's solid angle.

We verified the numerically computed focusing of the aerodynamic lens by exposing silicon targets to a ~90 nm average diameter polystyrene nanoparticle beam. Using a binocular loupe and image processing, the deposit diameter was estimated to be approximately 300 μ m (cf. Figure 5E). This value was reported on Figure 5F and is consistent with the numerical model predictions. It is worth noting that this represents a factor 5 improvement over the previous system generation (Dominguez-Medina et al., 2018) in terms of beam diameter. This is mainly due to the smaller lens-to-detector distance which used in the novel design (4 ν s. 8 cm). According to these results, as for the NEMS sensor, the aerodynamic lens may be used without performance degradation over the studied pressures for the size range of



interest, confirming that the NEMS-MS architecture is compatible with a substantial reduction of the pumping system.

3.3 Compact and modular NEMS-MS design

Following the aerodynamic lens study and the NEMS performance analysis over a wide pressure range, the vacuum system was modelled as described in Section 2.4 and a new pumping strategy was defined to maintain a 0.1 slm mass flow and related pressures of 2.7 mbar downstream the lens and 1.3×10^{-2} mbar inside the sensor chamber. Instead of the two turbomolecular pumps (450 l/s and 250 l/s) used to pump the system downstream of the aerodynamic lens and in the sensor chamber respectively, a single smaller turbo-molecular pump (TwisTorr 74FS, Agilent Technologies, Les Ullis, and France) was used to pump down the sensor chamber. A backing primary scroll pump (IDP-3, Agilent Technologies, Les Ullis, and France) was also used to pump the chamber downstream of the aerodynamic lens and back up the turbo-molecular pump.

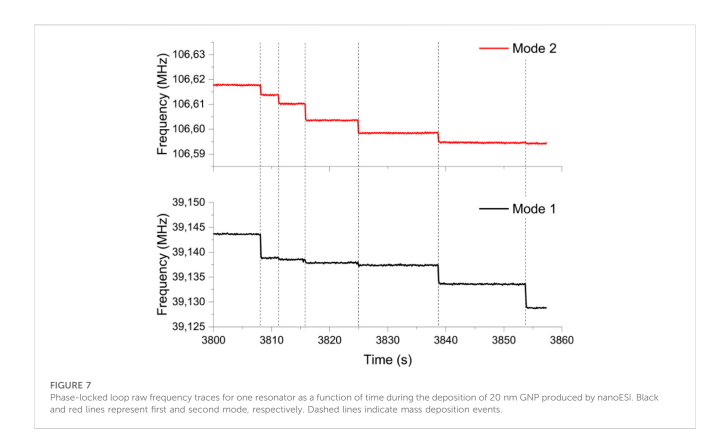
One of the objectives underlying the reduced pumping is to downsize the NEMS-MS bench in order to make it more practical to use. Thus, this second-generation prototype was engineered considering not only the dimensions, but also the modularity of the instrument. This will allow future users facing novel challenges to upgrade the instrument. Therefore, each function (particle intake, aerosol focusing and particle mass measurement) has been attributed a different mechanical module, as shown on Figure 6. Having opted to limit the diameter of the instrument, the mechanical cohesion of the assembly could be ensured by KF50 flanges, which are easier to mount, open and close. Moreover, the overall system was mounted on a rail, keeping the sensor chamber fixed and allowing the inlet chamber and the aerodynamic lens to slide away from the nano-resonator housing to access and replace the sensor's chip.

Because of the heavy pumps operated on the first-generation prototype, heavy vacuum components were used to support the weight and to prevent mechanical vibrations. Large hardware parts made the aerodynamic lens-to-detector distance optimization difficult and constrained the sensing area to deposit area ratio. Moreover, large mechanical parts being complex and expensive to machine, a built-in alignment solution could not be implemented, and a port aligner as well as a XY stage were used to align the lens outlet and the NEMS array respectively. As a result, one of the most time-consuming tasks to perform when replacing a NEMS chip was the alignment process, which was required to keep the sensor at the center of the particle beam produced by the aerodynamic lens. This operation was facilitated in the new system by designing a native alignment between the inlet capillary, the aerodynamic lens housing and the skimmer. Henceforth, the alignment step reduces to positioning the sensor within the particle beam area instead of aligning multiple parts aerodynamic lens and sensor as in the previous NEMS-MS prototype.

Finally, since the novel NEMS-MS prototype may host various sensor technologies—Nano-electromechanical as well as nano-optomechanical devices—The sensor chamber was designed accordingly, guaranteeing an easy and compact alignment solution whatever the type of chip used. The sensor itself is packaged onto a printed circuit board which is mounted on a motorized XY stage (Agilis AG-LS25, Newport, Irvine, California, United States), which could be used to monitor the sensor location and scan the particle beam.

3.4 Gold nano particles mass spectra

Eventually, to assess the new-generation compact and modular NEMS-MS system, we performed mass analysis of a series of calibrated samples of GNP of sizes 20, 30, and 40 nm. Nanoparticle colloidal suspensions were diluted in methanol, achieving concentrations of 4.50×10^{10} to 3.50×10^{11} particle/ml depending on nanoparticle diameter. Figure 7 presents examples of frequency traces obtained in both resonance modes during exposition to 20 nm GNP. The amplitudes of the observed frequency discontinuities can be translated into the mass-position domain, allowing the production of a particle mass distribution (PMD) (cf. Section 2.2.1). The distributions obtained for the three sampled analyzed in this study are shown in Figure 8.



Since GNP manufacturers characterized the batches by measuring nanoparticle diameters, the PMD were translated into mass equivalent diameters, assuming perfect sphericity and homogeneous density equal to that of bulk gold $(\rho_{gold} = 19.3 \, g/cm^3)$. Gaussian functions were used to fit masses and diameters histograms. Fitting parameters are reported in Table 2 and show good agreement with manufacturer's data (diameter measured by SEM), the relative difference in size ranging from 0.3% to 16% depending on the sample, and from 8.0% to 11.9% in mass. The standard deviation also compared well with manufacturer's data. Moreover, it should be highlighted that even if this comparison allowed to quantify the difference between NEMS-MS mass measurement and SEM size measurement, these techniques do not determine the same metric. Indeed, in order to compute the diameter of a particle, SEM is based on 2D image processing and NEMS-MS performs a mass-to-size conversion. Thus, both techniques assume the sphericity of particles, but the different metrics may induce different error factors. Finally, one might keep in mind that nano-resonator mass measurement has its own uncertainty as investigated previously by Clement et al. (2021).

In addition to characterizing mass measurement accuracy and precision, these experiments allowed to estimate the particle capture efficiency, which is useful for comparison with the previous system. Particle capture efficiency, as defined as the ratio between the number of detected particles to the number of GNP nebulized from solution during the measurement, was determined for one single NEMS among the 20 resonators of the array. The measurement details (i.e., flow rate, duration, initial concentration) are presented in the Supplementary Material

(Section 8). The particle capture efficiencies were computed for each GNP sample and are reported in Table 3. These values are compared with those reported by Dominguez-Medina et al. (2018), performed with ESI nebulization technique, showing an improvement factor of 20-200. The detection efficiency improvement could partly be attributed to the reduction in aerodynamic lens-to-sensor distance which has been halved (4 cm vs. 8 cm). Another possible reason may be that the new prototype is less affected by pump vibrations as the lens housing and the skimmer are mounted contiguously. Moreover, the lens housing is also aligned with the rest of the mechanical parts of the system, and has no degrees of freedom. These mechanical optimizations indirectly benefited from the pumping requirements downsizing, which allowed to reduce every dimension of the system, and were not possible on the firstgeneration prototype due to the cumbersome vacuum hardware components.

The interest in analyzing large sample fractions grows when the amount of available sample volume is small, which is often the case for biological samples, or for broad mass distributions that require a lot of events to acquire adequate statistics. To pursue the same objective (Erdogan et al., 2022), chose to dispense with vacuum system altogether and performed on-chip electrostatic focusing at ambient pressure. The main benefit of this technique was to reduce the volume and the cost of the apparatus while achieving a particle capture efficiency of 1 per 1.85×10^5 particle to 1 per 4.97×10^5 particle for one device. At this point, it should be stressed that particle capture efficiency comparisons are complicated due to potentially different raw resonance frequency postprocesses.

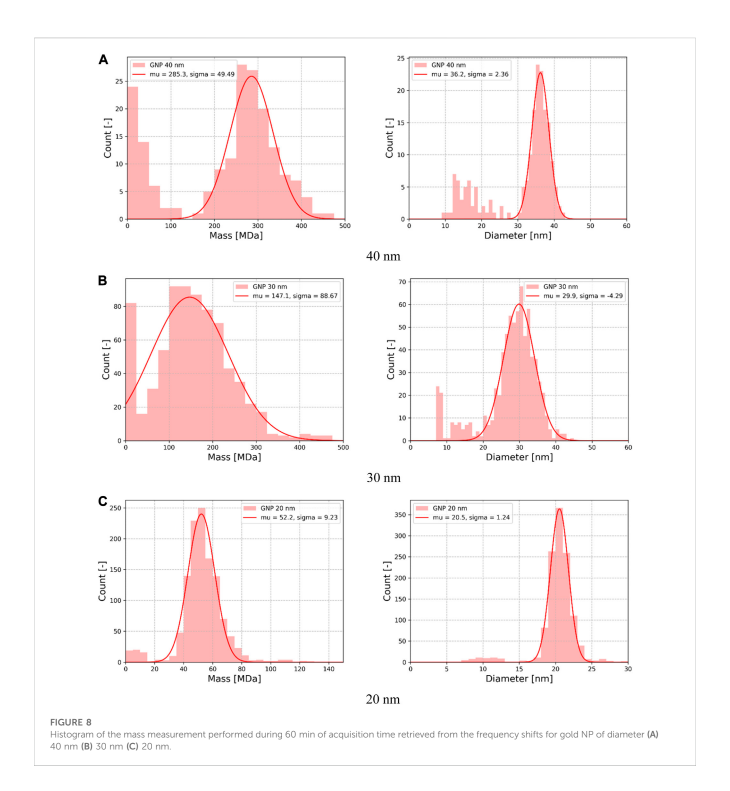


TABLE 2 Gaussian fit parameters (mean and dispersion) compared with manufacturer data.

		Mass		Diameter		
Sample	NEMS-MS $ar{m{m}}$ ($\sigma_{m{m}}$)	SEM m	Relative difference	NEMS-MS $\overline{m{d}_{p}}$ $(\sigma_{m{d}_{p}})$	SEM $\overline{d_p}$	Relative difference
	[MDa]	[MDa]		[nm]	[nm]	[-]
20 nm (BBI)	52.2 (9.2)	46	11.9%	20.5 (1.2)	19.6 (<1.6)	4.4%
30 nm (Sigma-Aldrich)	147.1 (88.7)	164	11.5%	29.9 (4.3)	30 (2)	0.3%
40 nm (BBI)	285.3 (49.5)	308	8.0%	36.2 (2.4)	37-42 (N/A)	2.2%-16%

TABLE 3 Capture efficiency computed for the gold nanoparticles. ¹Range computed based on the upper and lower capture efficiency reported by (Dominguez-Medina et al., 2018), namely 1 per ~2×10⁹ and 1 per ~1×10¹⁰ sprayed particle, respectively. ² Capture efficiency expressed as 1 detected particle per N nebulized particles.

GNP sample	Sprayed particles	Number of NEMS	Event count	Capture efficiency ²	Capture efficiency for 1 NEMS ²	Improvement factor ¹
	[× 10 ⁹ NP]	[-]	[-]	[-]	[-]	[-]
20 nm (BBI)	12.3	9	1220	1 per 4.5 × 10 ⁶	1 per 9.0 × 10 ⁷	22-111
30 nm (Sigma- Aldrich)	3.12	12	739	1 per 2.5 × 10 ⁶	1 per 5.1×10^7	40-198
40 nm (BBI)	1.58	9	183	1 per 3.9 × 10 ⁶	1 per 7.8 × 10 ⁷	26-129

4 Conclusion

In order to downsize the first generation NEMS-MS prototype without compromising mass measurement quality, the impact of an increase in operating pressure was analyzed. For this purpose, we separately addressed the NEMS and the aerodynamic lens. NEMS devices were characterized experimentally over a wide pressure range $(10^{-3}-10^3 \text{ mbar})$ and it was shown that their performance was unaffected at pressures up to 0.5 mbar. The aerodynamic lens was simulated numerically over an operating downstream pressure range of $1.3 \times 10^{-2}-1.3$ mbar for particle within the range of 25–500 nm and did not show to be significantly affected by this parameter.

Based on these results, a new vacuum system has been designed with a single turbomolecular pump (74 L/s), replacing the two larger pumps used in the previous-generation assembly (450 L/s and 250 L/s). This radical optimization led to a more compact, lightweight, modular and easy to operate mechanical design. Moreover, it allowed to position the NEMS sensor closer to the aerodynamic lens outlet, improving the particle transfer from the source to the sensor. Performance improvement was quantified using two approaches: the particle beam area at the sensor has been measured to be approximately 5 times smaller than in the previous design, and the particle capture efficiency has been increased by a factor of 20–200. The prototype, based on arrays of NEMS operated with a frequency-addressing scheme, was validated with mass measurements of standard nanoparticle samples and showed excellent agreement with predicted values.

Particle capture efficiency being one of the main limitations of NEMS-based MS, it was one of the main motivations underlying this design optimization. Improving it allows to perform analysis faster: the data used to plot the spectra reported on Figure 8 were registered during 60 min experiments and would have required more than 10 hours on the previous setup. Overall, the NEMS-MS technique presents a trade-off between particle capture efficiency and mass resolution: increasing the NEMS exposition to aerosol seems to imply an increase in pressure, which in return affects the mass resolution. In our system, the trade-off to keep a 0.1 MDa mass resolution (Clement et al., 2021) seems to be, pressure-wise, at 0.5 mbar (cf. Figure 3). However, this threshold might not be suitable for smaller particles as the diffusion could cause the divergence of the particle beam downstream the aerodynamic lens (Wang and McMurry, 2006).

Although the presented NEMS-MS prototype yielded significantly enhanced performance, there is still potential for

improvement. For instance, the size of the aerodynamic lens can be decreased, as it is the longest component in this prototype. Moreover, due to pumping considerations, the actual NEMS operating pressure $(1 \times 10^{-2} \text{ mbar})$ is still lower than the identified limit of operation (5×10^{-1} mbar), which opens the way to further developments using even smaller pumps or inlets with larger intake. Furthermore, detection efficiency could be further increased by NEMS and aerodynamic lens architecture, while sensitivity and mass range could be enhanced by NEMS sensor geometry. For the latter purpose, nano-optomechanical sensor might be implemented in this apparatus, which offer larger sensitive area and mass determination independently of particle landing position and physical properties (stiffness, size, and shape) (Sansa et al., 2020). These optimizations will eventually lead to new iterations of this NEMS-MS prototype, which will be greatly facilitated by its modularity.

Data availability statement

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

Author contributions

AR, CM, and SH conceived and designed the novel instrument. AR, LD, VÇ, and WT built and tested the instrument. AR, CM, MS, SH, and WT conceived and designed the simulations and experiments. AR, LD, VÇ, and WT performed the simulations and experiments. AR, CM, LD, MS, TF, and WT processed, analyzed, and interpreted the data. AR, CM, and WT drafted the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest

CM and SH are co-inventors of patents US9506852B2, EP2779209A1, and JP6352004B2.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2023.1238674/full#supplementary-material

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Effect of sub-micron grains and defect-dipole interactions on dielectric properties of iron, cobalt, and copper doped barium titanate ceramics

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Introduction: Dilutely doped ferroelectric materials are of interest, as engineering these materials by introducing point defects via doping often leads to unique behavior not otherwise achievable in the undoped material. For example, B-site doping with transition metals in barium titanate (BaTiO₃, or BTO) creates defect dipoles via oxygen vacancies leading enhanced polarization, strain, and the ability to tune dielectric properties. Though defect dipoles should lead to dielectric property enhancements, the effect of grain size in polycrystalline ferroelectrics such as BTO plays a significant role in those properties as well.

Methods: Herein, doped BTO with 1.0% copper (Cu), iron (Fe), or cobalt (Co) was synthesized using traditional solid-state processing to observe the contribution of both defect-dipole formation and grain size on the ferroelectric and dielectric properties.

Results and discussion: 1.0% Cu doped BTO showed the highest polarization and strain ($9.3\,\mu\text{C/cm}^2$ and 0.1%, respectively) of the three doped BTO samples. While some results, such as the aforementioned electrical properties of the 1.0% Cu doped BTO can be explained by the strong chemical driving force of the Cu atoms to form defect dipoles with oxygen vacancies and copper's consistent +2 valency leading to stable defect-dipole formation (versus the readily mixed valency states of Fe and Co at +2/+3), other properties cannot. For instance, all three T_c values should fall below that of undoped BTO (typically $120^{\circ}\text{C}-135^{\circ}\text{C}$), but the T_c of 1.0% Cu BTO actually exceeds that range (139.4°C). Data presented on the average grain size and distribution of grain sizes provides insight allowing us to decouple the effect of defect dipoles and the effect of grain size on properties such as T_c , where the 1.0% Cu BTO was shown to possess the largest overall grains, leading to its increase in T_c .

Conclusion/future work: Overall, the 1% Cu BTO possessed the highest polarization, strain, and $T_{\rm c}$ and is a promising dopant for engineering the performance of the material. This work emphasizes the challenge of extricating one effect (such as defect-dipole formation) from another (grain size modification) inherent to doping polycrystalline BTO.

KEYWORDS

barium titanate, doping, ferroelectric, grain size, ceramic processing

1 Introduction

Engineering ferroelectric ceramics by purposefully introducing point defects has a profound effect on the material's behavior, such as unusual ferroelectric properties and high electric field induced strain (Zhao et al., 2019). Examples of point defects include dopant or impurity ions as well as vacancies, which can lead to defect dipoles arising from oxygen vacancies and ionized acceptors (Ren, 2004; Maier et al., 2015). For perovskites (ABO₃ structure), dilute doping at the B site leads to a distribution of oxygen vacancies to achieve charge neutrality, which form defect complexes with the doped acceptor ion (Vani and Kumar, 2012; Maier et al., 2015). The oxygen vacancies have an intrinsic mobility, which is affected by the dopant itself, the concentration of dopant, and the transition metal's valence state. Barium titanate (BaTiO₃, BTO) is a ubiquitous ferroelectric oxide with low dielectric loss, chemical stability, and is lead-free with low toxicity, and has been widely and commonly doped at the Ti⁴⁺ site with various transition metals, including iron (Fe), manganese (Mn), cobalt (Co), and copper (Cu) (Rani et al., 2016). Although there are a multitude of studies on such dopants, these works focus on dopant concentrations that lead to hexagonal BTO, do not include polarization/strain curves in addition to permittivity and loss measurements, study ferromagnetic or optical properties, are processed under different thermal profiles, and do not attempt to directly compare three different dopants in the same work (Langhammer et al., 2003; Cheng et al., 2005; Dang et al., 2011; Deka et al., 2014; Langhammer et al., 2015; Langhammer et al., 2016; Rani et al., 2016; Krimech and Sayouri, 2020; Althobaiti, 2023).

In perovskites, such as dilutely doped BTO, both the ionic radius and valency of the dopant is highly important in determining the site of doping (either A or B site). This is closely related to how effective the particular dopant will be at trapping oxygen vacancies, which is highly influenced by the valence state and the charge neutrality conditions of the composition (Eichel, 2008; Schie et al., 2014). In addition, the concentration of oxygen vacancies is influenced by the concentration and specific element chosen as the dopant (Huang et al., 2014). B-site dopants with a lower valency than Ti⁴⁺ (i.e. 2+ or 3+) can then form stable defect dipoles with oxygen vacancies along the <001> direction, the same direction as the spontaneous polarization direction (Zhao et al., 2019). These particular defectdipole complexes have been shown to have a profound effect on the ferroelectric and piezoelectric behavior of materials such as barium titanate. The moment of the defect dipole induces an internal bias field, which can increase the polarization of the material by producing large crystal field anisotropy in the vicinity of the dopant and, when poled in the direction of the defect dipole, can also increase the strain experienced in the material (Maier et al., 2015). The orientation of the dipoles takes place during oxygen vacancy diffusion on the scale of several unit cells, which occurs at room temperature and leads to a recoverable domain wall configuration and "pinched" hysteresis loops with decreased remnant polarization (Kamel and de With, 2008; Zhao et al., 2019). Overall, a dopant cation has a direct effect over the range of several unit cells on the association and migration energies, i.e., the energies of oxygen vacancy-dopant association and activation energy of vacancy motion away from the dopant, respectively (Schie et al., 2014). This can significantly impact the formation of these complexes and thus the bulk ferroelectric properties.

Although our previous work (Patterson et al., 2023) and several other works discussed above focus on single crystals, for polycrystalline samples another important influence on the ferroelectric and piezoelectric properties of both doped and undoped BTO is grain size. It has been reported that several electrical properties such as the piezoelectric constant (d₃₃), polarization, and relative permittivity are affected by grain size, typically with a decrease in these properties with a decrease in grain size as grains approach the nanometer scale (Zhao et al., 2004; Guo et al., 2012; Huan et al., 2014). This is predicted to be due to a change in domain density and non-uniformities at grain boundaries (Huan et al., 2014). It has also been observed that non-uniformities such as aluminum segregation at grain boundaries can be partially responsible for ferroelectric aging behavior even when no second phase is detected, separate from impacts of volume of domain walls (Carl and Hardtl, 1977). Overall, domain configuration has a large impact on macroscopic dielectric properties, which in turn is substantially affected by the microstructure (Eichel, 2008). However, it is imperative to attempt to separate the effects of microstructure and intrinsic effects on the dielectric properties of polycrystalline ferroelectrics.

In this work, dielectric materials properties will be investigated (polarization, strain, and relative permittivity/loss) and reported with respect to polycrystalline BTO ceramics doped with 1.0 mol% of Fe, Co or Cu, synthesized via traditional solid-state processing. These results will then be discussed with respect to the grain size, determined from cross-sectional scanning electron microscopy (SEM) images. Comparisons between the dielectric properties of the three different doped BTO samples show the importance of considering all of the various contributions from the dopant that can affect the dielectric properties. These can be due to defect-dipole formation, grain growth/growth inhibition, oxygen vacancy diffusion, or a complex contribution from a subset or all of these. The challenge of extricating one effect from the other will be emphasized, along with the need for these in-depth studies.

2 Materials and methods

Ceramic powders of BaTi_{1-x}Me_xO₃ (with Me = Fe, Co, or Cu) were prepared using traditional solid state synthesis methods with x = 0.01 mol. Precursor oxides of BaCO₃, TiO₂, Fe₂O₃, CoCO₃, and CuO (99.9+%) were mixed in ethanol and ball milled for 8 h with proper stoichiometric amounts of the Fe, Co or Cu precursor added. The mixed powders were dried and then calcined at 1,150°C for 2 h and milled and dried again as before. Pellets of each composition were then uniaxially pressed in a 12.7 mm diameter die after mixing the calcined powder with 4-5 wt% binder (Paraloid B-72 resin mixed with acetone). The pressed pellets were sintered at 1,250°C in air for 4 h. The density of the sintered pellets was greater than 91% theoretical density for all samples in this study. The phase of the pellets was confirmed with X-ray diffraction (XRD) using a Rigaku SmartLab Rigaku X-ray powder diffractometer with Cu Ka radiation at 40 kV and 200 mA for a 20 range from 20° to 80° with a 0.02° step size.

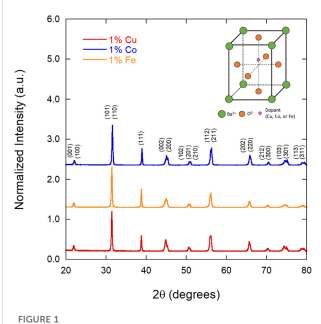


FIGURE 1
X-ray diffraction patterns of the 1% Fe, Co, and Cu-doped BTO powders measured after calcination at 1150°C, showing the tetragonal phase of BTO and no evidence of a hexagonal phase impurity (indexed with respect to PDF# 04-012-8129). The crystal structure of tetragonal BTO is shown in the top right corner.

The sintered pellets were then ground down to a thickness of approximately 800 µm and masked and sputtered with gold electrodes for synchronous ferroelectric and piezoelectric measurements, which were performed with a self-constructed Sawyer-Tower system and a linear variable differential transformer (LVDT) connected to a lock-in amplifier. The electrical measurements were performed on samples in three different states with increasing applied voltages from 0.1 kV up to 1.4 kV at 1 Hz (all data shown in this work is at 1.4 kV) as the properties are well-understood to strongly depend on the defect configuration, which can be modified through temperature or electric field. Therefore, the thermal and electrical history of the sample are important factors in discussing the dielectric behavior. The three states to be measured are referred hereafter as: as-sintered, aged, and poled. All samples were thermally reset at 300°C for 4 h after the initial sintering and allowed to cool to room temperature. This default "as-sintered" state was then tested. The "aged" subset of these samples were then aged at 80°C for 24 h to allow for diffusion of oxygen vacancies prior to characterization. Finally, the "poled" subset of samples were poled at a DC bias of 1.5 kV for 1 h at room temperature and tested.

Simultaneous capacitance and tan δ measurements were performed using two different methods in order to determine the relative permittivity and dielectric loss of the samples under different conditions. All samples for these measurements were tested in the "as-sintered state." First, an Agilent E4980A precision LCR meter was used to sweep between frequencies of 200 Hz and 2 MHz to determine the permittivity and loss at room temperature. Second, a Thermal Product Solutions (TPS) Tenney Environmental Test Chamber was used to measure the capacitance and loss from -30°C to 155°C at four different frequencies: 0.1, 1, 10, and

100 kHz. The frequency was generated from an HP 4284A Precision LCR meter, which also measured the capacitance and loss at each frequency over the temperature range. A self-assembled LCR test interface was used to connect the LCR meter to the sample holder inside the environmental test chamber, which was accompanied by a National Instruments thermocouple placed close to the sample to more accurately monitor temperature.

Cross-sectional scanning electron microscopy (SEM) images were taken of the polished, fractured surface of each composition using a Thermo Scientific Quattro S at an accelerating voltage of 10 keV. Images were taken across different regions of the cross-section, and were analyzed by ImageJ software. Ten images were used to determine the average grain size via the linear intercept method, whereby five lines were drawn per image to obtain an average grain size for one image, resulting in 50 total lines drawn with ten average values of grain size for each particular sample.

3 Results and discussion

3.1 X-ray diffraction

Sintered pellets of 1% Fe, Co, and Cu-doped BTO were analyzed via XRD to determine if the concentration of Fe affected the presence of the expected ferroelectric, tetragonal phase, as some studies have observed the formation of the hexagonal phase at as low as 1% Fe (Langhammer et al., 2020). Figure 1 shows that across the three samples, the calcined powders are of the ferroelectric tetragonal phase with no other phases present, demonstrating that doping in this study with Fe, Co, or Cu does not induce a detectable phase change at x = 1%. The lattice parameters and c/aratios obtained from a whole pattern fit of each diffraction pattern (indexed with respect to tetragonal BTO, PDF# 04-012-8129) is included in Table 1. The c/a ratio of undoped BTO, 1.01, is slightly higher than that of the Co, Fe, and Cu-doped BTO samples (1.008, 1.007, and 1.008, respectively). Doping with 1% Co and Fe has been shown to slightly decrease the c/a ratio, which was observed here for all three dopants (Buscaglia et al., 2000).

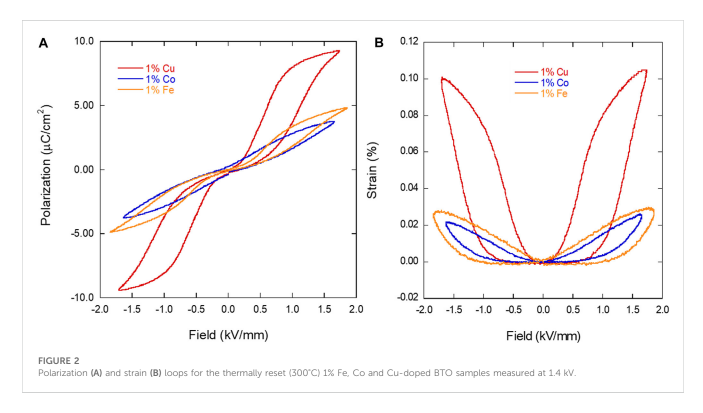
3.2 Ferroelectric and piezoelectric testing

Polarization and strain of the three compositions in all three conditions were measured at 1.4 kV, which translates to a field of approximately 1.5 kV/mm. Figure 2 shows polarization (Figure 2A) and strain loops (Figure 2B) for the "as-sintered" condition measured at 1.4 kV. Note that all samples were tested at applied voltages between 0.1 and 1.4 kV at 1 Hz, but only the 1.4 kV results are shown here. The polarization and strain loops for the samples that were aged (80°C, 24 h) and poled (1.5 kV, 1 h at 2.4 kV) can be found in the Supplementary Material (Supplementary Figures S1, S2). Measuring samples after aging and poling was performed to determine the extent that temperature and field cycling affects the polarization and strain of these differently doped materials. The 1.0% Fe-doped BTO sample showed an increase in strain from ~0.025% to ~0.04% after aging, and the 1.0% Cu BTO sample showed a decrease in polarization from ~9 to ~5 µC/cm² (Supplementary Figure S1). In order to fully realize the effect on

TABLE 1 Lattice parameters, dielectric properties, average grain sizes, and densities for the three doped BTO samples.

	Lattice parameters		@ 10 kHz						
	a,b	С	c/a		tan δ	T _{CH} (°C)	T_{c_C} (°C)	Avg. grain size (μm)	Relative density (%)
1% Co BTO	4.002	4.033	1.008	1830	0.017	107.1	90.2	0.62 ± 0.2	95.2
								2.29 ± 0.3	
10/ E PEO	4.005	4.022	1.007	22.40	0.012	00.0	(1.0	0.88 ± 0.2	0.0
1% Fe BTO	4.005	4.033	1.007	2240	0.012	99.8	61.2	49.19 ± 13.5	96.9
								4.73 ± 1.1	
1% Cu BTO	4.004	4.036	1.008	2150	0.026	139.4	103.6	14.35 ± 4.4	91.4
								32.57 ± 3.8	

Dielectric properties are reported for samples measured at 10 kHz, and the Curie temperature is reported both on heating (T_{c_0}) and on cooling (T_{c_c}) . Lattice parameters were obtained from a whole pattern fit of the XRD patterns of each sample. Grain size was measured using the linear-intercept method. Density was measured using the Archimedes method.



the polarization and strain after poling, the poled samples were measured at 2.4 kV (Supplementary Figure S2), showing multiple changes in polarization and strain behaviour. It is important to not compare their maximum values without considering that the field level is nearly twice as high as in the as-sintered and aged state. With that in mind, the 1.0% Cu BTO sample displayed no further enhancement compared to the "as-sintered" value of polarization and the pinched portion of the polarization hysteresis loop disappears in favor of a traditional ferroelectric shape. The strain also takes on a more butterfly shape with negative strain at the coercive field. Additionally, the polarization loops of the 1.0% Co and Fe samples have become asymmetric, indicating a high internal electric field (Supplementary Figure S2A). This is supported by the offset in the field of the 1.0% Co and Fe BTO strain loops (Supplementary Figure S2B). The poled strain loops also show a

general improvement of strain for all three samples that approaches 0.1%, albeit at higher fields as mentioned. Aging and poling can have extreme effects on the polarization and strain behaviour of ferroelectric materials via domain/defect-dipole orientations (Kamel and de With, 2008). While the effects that both aging or poling have on the polarization and strain of the material via an external, post-processing method are quite large, in these samples they were found to be repeatably reversible following a 300°C heat treatment. Importantly, the hysteresis loops always returned to the "as-sintered" state, so it is from this baseline condition that comparisons will be focused on in this discussion.

In the "as-sintered" state, the 1.0% Cu-doped BTO yields the highest polarization (~9.3 μ C/cm²) and strain (~0.1%) when compared to the 1.0% Fe and Co-doped BTO, with polarizations of 4.8 and 3.7 μ C/cm² and strains of 0.03% and 0.025%, respectively.

In comparison, undoped BTO sintered at similar conditions yield polarizations between 15 and $18\,\mu\text{C/cm}^2$ (Tan et al., 2015; Malyshkina et al., 2020). Additionally, Figure 2B shows that all three dopants yielded little to no negative strain, showing recoverable strain across each dopant. The presence of the pinched polarization loops of the 1.0% Fe and Cu-doped BTO samples (Figure 2A) is a good indication of defect dipole formation pinning domain walls in those samples.

There is an obvious improvement in the ferroelectric properties with respect to the 1.0% Cu-doped BTO when compared to the 1.0% Fe and Co. Looking at each dopant, however, it does not seem at first glance that ionic radius plays a part, as the difference between the ionic radii for the various oxidation states of each transition metal are relatively similar when compared to Ti⁴⁺ (0.61) (listed in Å): 0.75 (Co²⁺), 0.55 (Co³⁺), 0.77 (Cu⁺), 0.73 (Cu²⁺), 0.78 (Fe²⁺), and 0.65 (Fe³⁺) (Shannon, 1976). Therefore, doping with any of these three elements for Ti⁴⁺ should not play a role in one composition more than another. It is important to note, however, that the valence state of the incorporated ion is dependent on a number of factors, such as initial precursors and the oxidizing/reducing conditions during synthesis. Therefore, we should expect a mix of these valence states for some if not all of the dopants chosen in this work.

While the valence states of the incorporated dopants will not significantly distort the lattice, the valency of the ions can influence the intrinsic ferroelectric behaviour. Co and Fe are both stable at a 2+ or 3+ state, while Cu is commonly found in a 2+ state. The mixed valence of the Co and Fe may play a role in achieving charge neutrality (via different amounts of oxygen vacancies and other free charges) which then affects the ferroelectric and piezoelectric properties of the material. This difference in valence is proposed to directly affect the establishment of defect-dipoles, formed between the dopant and induced oxygen vacancies. Using Kröger-Vink notation, the defectdipole complexes for the three different dopants are the following: $(Fe'_{Ti}-V_o^*)^T$, $(Co'_{Ti}-V_o^*)^T$, and $(Cu''_{Ti}-V_o^*)^X$. The charge states of the three different defect dipole complexes are based on the assumption that Fe and Co are both in a 3+ state and Cu is 2+. This was assumed because it has been shown that, for Fe and Co, oxidation from the 2+ to 3+ state is more energetically favourable in comparison to the incorporation of those two elements in their divalent state (Buscaglia et al., 2001). Due to the charged nature of the defect-dipole complexes for Fe and Co in the 3+ state, there are additional mechanisms for charge compensation that may take place, including free electrons trapped in the lattice (Eichel, 2008). This excess charge can give rise to a screening effect that ultimately causes ferroelectric instability, which could explain the decreased polarization and strain in the 1% Fe and Co-BTO samples when compared to the 1% Cu sample (Wang et al., 2012). Additionally, work by Erhart et al. (2007) suggests that there is a very strong chemical driving force between copper dopants and oxygen vacancies in lead titanate, which theoretically leads to all copper ions complexed with an oxygen vacancy.

3.3 Other dielectric measurements

3.3.1 Relative permittivity (ϵ_r) /tan δ vs. frequency (room temperature)

The relative permittivity (ϵ_r) and loss tangent (tan $\delta)$ of assintered samples of each composition were measured at room

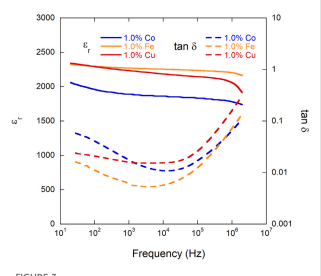


FIGURE 3 Relative permittivity (ϵ_r) (solid lines) and loss tangent (tan δ) (dashed lines) of each composition (1% Co, Cu, and Fe BTO) measured at room temperature with respect to frequency after samples were thermally reset at 300°C for 4 h.

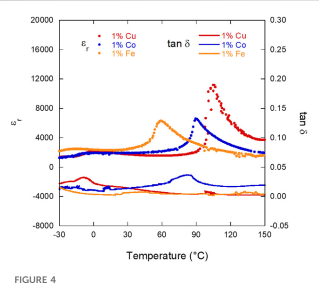


FIGURE 4 Relative permittivity (ϵ_r) (solid lines) and loss tangent (tan δ) (dashed lines) of each composition (1% Co, Cu, and Fe BTO) with respect to temperature, measured at 10 kHz in the "as-sintered" state (thermally reset at 300°C for 4 h after sintering). Only the cooling curves are shown in this plot.

temperature with respect to frequency, shown in Figure 3 and reported in Table 1. When compared to the typical relative permittivity of barium titanate (ϵ_r ~3,000-6,000), all three of the doped BTO samples exhibit a lower value than undoped BTO, which is due to the fact that in comparison to Ti⁴⁺, all three dopants (Co³⁺, Cu²⁺, and Fe³⁺) are more conductive and less polarizable, decreasing the relative permittivity, as expected (Rani et al., 2016). More specifically, undoped BTO at similar processing conditions yields a relative permittivity similar to the reported ranged of BTO, ranging

from ~3,000 to 6,500 at 1 kHz (Hoshina et al., 2008; Tan et al., 2015). However, the permittivity of all three doped BTO samples remains fairly constant up to 2 MHz, due to the failure of the dipoles to follow the fast alternating electric field (Rani et al., 2016). With regards to tan δ at 10 kHz, each doped sample exhibited a loss of ~0.01, a relatively low value of loss due to the dopant's ability to reduce leakage current in the sample (Imai et al., 2002). All samples showed an increase in loss in the MHz, frequency range and the 1% Cu doped BTO sample nominally yielded the highest loss at 2 MHz. The Cu doped sample, however, showed the most frequency-stable loss at the intermediate to lower frequencies (with the increase starting at 100 kHz).

3.3.2 Relative permittivity (ϵ_r) /tan δ vs. temperature (variable frequency)

As-sintered samples of 1% Co, Cu, and Fe doped BTO were loaded into an LCR test interface and the ε_r and tan δ were measured between -30°C and 155°C at 0.1, 1, 10, and 100 kHz. Data from the cooling curves for each sample at 10 kHz are shown in Figure 4 with Curie temperatures measured both on heating (T_{c_H}) and cooling (T_{c_c}) are reported in Table 1. Plots of ε_r and tan δ with respect to temperature at all four frequencies are included in the Supplementary Material (Supplementary Figure S3). The magnitude of ε_r is highest at the Curie temperature for 1% Cu $(\sim 7,200)$ and is similar for 1% Co and 1% Fe $(\sim 5,520)$. Additionally, the T_{c_H} of the 1.0% Cu-doped BTO (139.4°C) is higher than both the 1.0% Co (107.1°C) and 1.0% Fe (99.8°C) samples. One can also observe that there is a slight increase in tan δ at T_{c_H} for all three dopants, and that the loss is consistently highest in the 1% Co sample. To further explore the differences in T_c values of the three dopants, the difference between the T_c on heating and cooling (ΔT_{Ch-Cc}) have been calculated as 16.9, 38.6, and 35.8°C for Co, Fe, and Cu doped BTO, respectively. The smaller ΔT_{Ch-Cc} of the 1% Co and the similar $\Delta T_{\text{Ch-Cc}}$ values of the 1% Fe and Cu doped BTO samples could be due to the relative abilities of each dopant to trap vacancies in the sample, which would lead either more or less thermal energy needed to complete the phase transition between the tetragonal and cubic phases.

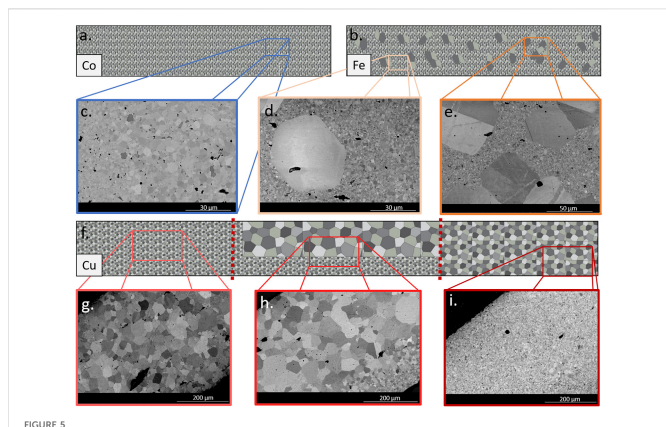
With respect to the T_{c_H} of the three samples, it is important to note that the accepted value of T_c for undoped BTO lies between 120°C and 135°C, with samples processed under comparable conditions yielding Curie temperatures ranging between 120°C and 130°C (Tan et al., 2015; Malyshkina et al., 2020). It is generally expected that doping BTO with these cations or ones of similar valency will decrease the T_c due to oxygen vacancy formation (Tewatia et al., 2021). The 1.0% Co and 1.0% Fe samples follow this trend in lower T_{c_H} than the reported value of undoped BTO: 1.0% Co (107.1°C) and 1.0% Fe (99.8°C). The similarity in T_{c_H} of the 1.0% Co and Fe samples follows the similarity of their trends in other properties, such as polarization and strain, which could be due to their similarities in valence (3+), which would lead to similar defectdipole formation and subsequent effects on ferroelectric properties as a result. In contrast, the T_{CH} of 1.0% Cu BTO was 139.4°C, higher than the maximum of the accepted range of T_c of undoped BTO. This suggests that there may be another factor at play that is driving the T_{c_H} higher, as the introduction of oxygen vacancies should decrease the T_c. One such factor is the grain size in the material. Properties such as polarization and relative permittivity are affected by grain size, with a decrease in these properties as grains approach the nanometer scale.

3.4 Grain size measurement

In order to measure the grain sizes of the three samples, cross-sections of a pellet of each dopant were polished and then imaged via SEM. Grain size was then determined by the linear intercept method, utilizing ten SEM images to obtain average grain sizes for the different grain size distributions within each sample. Schematics of each sample illustrating the grain structure of each sample can be seen in Figures 5A, B, F for Co, Fe and Cu dopants, respectively. Micrographs from different sub-sections of these representative cross-section schematics are also included in Figures 5C–E, G–I. The average grain size values can be found in Table 1.

The cross-section of the 1.0% Co sample (Figures 5A, C) had a minor but distinctly bimodal distribution of grains (0.62 \pm 0.2 and $2.29 \pm 0.3 \mu m$), revealing a population of submicron grains. These two groups of grains were equally distributed throughout the cross-section of the sample, as illustrated in Figure 5A. Next, the 1.0% Fe sample (schematically shown in Figure 5B) also had a bimodal distribution of grains (0.88 \pm 0.2 and 49.19 \pm 13.5 µm), but here the subset of larger grains were substantially bigger compared to the Co sample. For clarity, it is important to note that the larger grains of this sample were measured individually (measuring 175 grains to obtain the average), while the smaller grains were measured by the linear intercept method. Unlike 1.0% Co BTO, 1.0% Fe BTO had clusters of these large grains surrounded by the smaller, submicron grains, shown in the SEM images in Figures 5D, E, which are representative of what was observed along the entirety of the cross-section. Finally, the 1.0% Cu sample (illustrated in Figure 5F) had three distinct areas along the cross-section (Figures 5G-I) and three different populations of grain sizes (4.73 \pm 1.1, 14.35 \pm 4.4, and 32.57 \pm 3.8 μ m). Overall, 1.0% Cu had the largest grains of the three different dopants, albeit not distributed evenly over the cross-section. Upon electron dispersive spectroscopy (EDS) analysis, there was a lack of apparent dopant segregation into grain boundaries or secondary phases in the cross-sections of the doped samples. Undoped BTO sintered under similar conditions have grains averaging approximately 4 µm in diameter, the size of which is very dependent on the sintering temperature and time (Hoshina et al., 2008; Tan et al., 2015; Malyshkina et al., 2020). The choice of dopant itself is likely driving the observed differences in grain sizes between the three samples, since they were sintered intentionally under the same conditions to achieve similar high density (>91%) rather than optimizing the sintering conditions to create a specific grain size. In previous doping studies, for instance, Co has been used to control grain sizes in ceramics to ${\sim}1~\mu m$ by inhibiting grain growth, and Cu was added as a sintering aid to encourage grain growth (Cheng et al., 2005; Mahmoudi et al., 2019).

This grain size analysis, then, can potentially inform some of the differences in dielectric properties between the three samples. Although doping should theoretically decrease the Curie temperature, this was not observed in the 1.0% Cu sample. This



Schematics illustrating the grain structure of cross sections of 1.0% Co BTO (A), 1.0% Fe BTO (B) and 1.0% Cu BTO (F) samples. Magnified areas are SEM images from each sample [1.0% Co (C), 1.0% Fe (D,E), and 1.0% Cu (G-I)] showing the grain structure and size distribution along the length of the cross section.

increase in T_c could then be contributed, in part, to the larger average grain size, as it has been shown that T_c increases as grain size increases (Tewatia et al., 2021). Additionally, although there are larger grains present in 1.0% Fe BTO, the dominant small grains of this sample and that of the 1.0% Co sample could explain the lower T_c, polarization, and strain values when compared to 1.0% Cu BTO. The permittivity values do not follow this trend, as the 1.0% Cu and Fe samples have similar values, while the 1.0% Co BTO possesses the lowest $\epsilon_{\rm r}$. However, since these values are not vastly different (Table 1) and do fall below that of undoped BTO (ranging from 3,000 to 6,000), it may not be a significant indicator of the dependence of this value on either dopant or grain size alone. This analysis, though, does highlight the importance of considering both the defect dipole and grain size effects on dielectric properties. Although some dopants are typically classified as more or less effective than others, such as if a dopant's valency and/or quantity added should lead to positive enhancements via defect dipole formation, the effect of grain size can outweigh the benefit of the defect dipoles with respect to dielectric properties.

4 Conclusion

In this work, we have studied the dielectric properties of polycrystalline B-site doped BTO with 1.0% copper (Cu), iron (Fe), and cobalt (Co), synthesized using traditional solid-state

processing, to observe the contribution of both defect-dipole formation and grain size on the ferroelectric and dielectric properties. 1.0% Cu doped BTO showed the highest polarization and strain (9.3 µC/cm² and 0.1%, respectively) of the three dopants in and a very clearly pinched hysteresis and recoverable strain behavior in the as-sintered state, with some changes in polarization and strain to all three doped samples after aging or poling. Additionally, 1.0% Cu BTO showed the highest Curie temperature (Tc) (139.4°C). With respect to defect dipole formation, these results suggested that the strong chemical driving force of the Cu atoms to form defect dipoles with charge neutrality induced oxygen vacancies leads to improved polarization and strain. However, since this would not explain the high Tc of the 1.0% Cu; the average grain size and distribution of grain sizes provided insight for this parameter instead. The 1.0% Cu BTO possessed the largest overall grains, which leads to an increased Tc. Grain size cannot be the only contributor to the improvement of properties, though, as the larger grain size would not account for the type of pinching in the polarization loop and increased strain, which we attribute to defect-dipole formation. Overall, this work emphasizes the challenge attributing a singular contribution, such as defect dipole formation via doping or grain size to dielectric properties of polycrystalline, doped BTO. Instead, it is critical to obtain a more comprehensive view of the material as a whole in order to inform causation and conclusions.

Data availability statement

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

Author contributions

SM carried out the majority of the experiments outlined in the manuscript, EP also contributed to the experimental efforts. SM, EP, and MS conceived of the ideas, and SM and MS were in charge of overall planning. SM drafted the manuscript with support from EP and MS. All authors contributed to the article and approved the submitted version.

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Supplementary material

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Can a single ammonia and water molecule enhance the formation of methanimine under tropospheric conditions?: kinetics of $\cdot CH_2NH_2 + O_2 (+NH_3/H_2O)$

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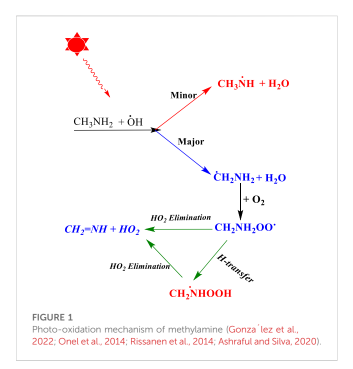
The aminomethyl (•CH₂NH₂) radical is generated from the photo-oxidation of methylamine in the troposphere and is an important precursor for new particle formation. The effect of ammonia and water on the gas-phase formation of methanimine (CH₂NH) from the ${}^{\bullet}$ CH₂NH₂ + O₂ reaction is not known. Therefore, in this study, the potential energy surfaces for *CH₂NH₂ + O₂ (+NH₃/H₂O) were constructed using ab initio//DFT, i.e., coupled-cluster theory (CCSD(T))//hybriddensity functional theory, i.e., M06-2X with the 6-311++G (3df, 3pd) basis set. The Rice-Ramsperger-Kassel-Marcus (RRKM)/master equation (ME) simulation with Eckart's asymmetric tunneling was used to calculate the rate coefficients and branching fractions relevant to the troposphere. The results show 40% formation of CH₂NH at the low-pressure (<1 bar) and 100% formation of CH₂NH₂OO[•] at the high-pressure limit (HPL) condition. When an ammonia molecule is introduced into the reaction, there is a slight increase in the formation of CH₂NH; however, when a water molecule is introduced into the reaction, the increase in the formation of CH₂NH was from 40% to ~80%. The calculated rate coefficient for ${}^{\bullet}\text{CH}_2\text{NH}_2 + \text{O}_2 \text{ (+NH}_3) [1.9 \times 10^{-23} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}] \text{ and for CH}_2\text{NH}_2 + \text{O}_2$ (+ H_2O) [3.3 \times 10⁻¹⁷ cm³ molecule⁻¹ s⁻¹] is at least twelve and six order magnitudes smaller than those for free ${}^{\bullet}\text{CH}_2\text{NH}_2 + \text{O}_2$ (2 × $10^{-11}\,\text{cm}^3$ molecule $^{-1}\,\text{s}^{-1}$ at 298 K) reactions, respectively. Our result is consistent with that of previous experimental and theoretical analysis and in good agreement with its isoelectronic analogous reaction. The work also provides a clear understanding of the formation of tropospheric carcinogenic compounds, i.e., hydrogen cyanide (HCN).

KEYWORDS

aminomethyl radical, O_2 radical, methanimine, ab initio/DFT, RRKM/ME, H_2O and NH_3 , HCN, catalysis

1 Introduction

Methylamine is a simple organic nitrogen compound that is released into the atmosphere from a range of sources, for example, food industries, animal husbandry, marine sources, and biomass burning (Schade and Crutzen, 1995; Ge et al., 2011a; Ge et al., 2011b; Zhang et al., 2012; Almeida et al., 2013). Methylamine forms a particulate salt when reacting with acids such as H₂SO₄, HNO, and CH₃COOH; therefore, it plays a vital role in enhancing



atmospheric cloud nucleation (Murphy et al., 2007; Lee and Wexler, 2013). The reaction of methylamine with various tropospheric oxidants such as O3, OH, and NO3 radicals leads to the formation of semi-volatile and non-volatile chemical species, consequently leading to the formation of secondary organic aerosols (Schade and Crutzen, 1995; Murphy et al., 2007; Ge et al., 2011a; Nielsen et al., 2012; Qiu and Zhang, 2013). Methylamine is also expected to be present in the interstellar medium (ISM), which leads to the formation of amino acids (Altwegg et al., 2016; Elsila et al., 2009; Gonza'lez et al., 2022). Although glycine (HO₂CCH₂NH₂) has not yet been identified in the ISM medium, it is detected in different comets (Elsila et al., 2009; Altwegg et al., 2016). Methylamines are also possible atmospheric precursors of hydrogen cyanide and nitrous oxide (N2O) (Nielsen et al., 2012). N2O is a greenhouse gas and the potential source of stratospheric NOx production. To know the significance of methylamine reactions in the two drastically environments, several researchers have investigated their atmospheric significance and sinks in both the gas phase and solid phase (Schade and Crutzen, 1995; Ge et al., 2011a; Ge et al., 2011b; Almeida et al., 2013).

Once CH₃NH₂ is released into the Earth's atmosphere, it reacts with the OH radical via the H-abstraction reaction, leading to the formation of a carbon-centered aminomethyl (${}^{\bullet}$ CH₂NH₂) radical, which is observed as a major product, and nitrogen-centered methyl amino radical (CH₃NH ${}^{\bullet}$), which is observed as a minor product (Figure 1) (Gonza'lez et al., 2022; Onel et al., 2014; Rissanen et al., 2014; Ashraful and Silva, 2020).

As suggested in the previous studies (Gonza'lez et al., 2022; Onel et al., 2014; Rissanen et al., 2014; Ashraful and Silva, 2020), •CH₂NH₂ predominantly reacts with molecular oxygen (O₂), which can lead to the formation of methanimine (CH₂NH) and the hydroperoxy radical (HO₂) as major reaction products via a hydrogen atom transfer (HAT) mechanism (Gonza'lez et al., 2022; Onel et al., 2014; Rissanen et al., 2014; Ashraful and Silva, 2020). The chemical kinetics studies on the CH2NH2 + O2 reaction system have been investigated by various researcher groups (Masaki et al., 1995; Jansen et al., 1999; Rissanen et al., 2014; Ashraful and Silva, 2020; Glarborg et al., 2020). Jansen et al. (1999) used pulse radiolysis and UV-absorption detection to analyze the chemical kinetics of the $^{\bullet}$ CH₂NH₂ + O₂ reaction at 298 K and 1 atm of SF₆ as a bath gas. Masaki et al. (1995) investigated the kinetics of the same reaction by employing the photoionization mass spectrometry technique at 298 K and a few torr pressure of N2. Rissanen et al. (2014) used the laser flash photolysis technique in combination with photoionization mass spectrometry to determine the rate coefficients of the CH2NH2 + O2 reaction. They observed negative temperature-dependent rate coefficients from 267 K to 363 K, independent of the pressure between 0.5 Torr and 2.5 Torr (Rissanen et al., 2014). The reported rate coefficients fall in the range of $(2-8) \times 10^{-11}$ cm³ molecule⁻¹ s⁻¹ (Rissanen et al., 2014; Mallick et al., 2018; Ashraful and Silva, 2020; Kumar et al., 2020). Rissanen et al. (2014) also performed quantum chemical calculations coupled with ME simulation to predict the product branching fractions. Their modeling results reproduce the experimentally observed negative temperature dependence and validated the production of CH2NH under low-pressure conditions. Recently, a chemical kinetic model for the oxidation of methylamine has been characterized by Glarborg et al. (2020) and validated against the results obtained from shock tube experiments. In their work, the potential energy surface of several reactions was studied theoretically starting from the isomerization of CH₃NH and the reactions of CNHx (x = 3-5) molecules with O₂ using the quantum chemistry composite method. In this study, the rate coefficients for CH2NH2 + O2 were re-investigated at high-level quantum chemical calculations with similar statistical rate theories to validate the finding for the role of ammonia and water and molecules in the same reaction. To the best of our knowledge, the branching ratios and temperature- and pressure-dependent rate coefficients have not been available until now.

Concerning the gas-phase reactivity of CH2NH2 toward O2 in the role of ammonia and water, several studies in the past few years have proposed the role of different species such as H₂O, NH₃, formic acid, and CO2 on important atmospheric reactions (Vöhringer-Martinez et al., 2007; Iuga et al., 2010; Buszek et al., 2012; Iuga et al., 2011; Thomsen et al., 2012; Jonas et al., 2013; Zhang et al., 2013; Zhang et al., 2014; Jara-Toro et al., 2017; Ali et al., 2018; Inaba, 2018; Mallick et al., 2018; Ali, 2019; Ali et al., 2019; Ali, 2020; Kumar et al., 2020; Wu et al., 2020; Zhang et al., 2020; Ali et al., 2021; Ali and Balaganesh, 2022; Ali et al., 2022; Dash and Ali, 2022). It is wellknown that ammonia (NH₃) is highly alkaline and is one of the most common chemicals used in the agriculture sector, and as a fertilizer, it is the major source in the atmosphere. Ammonia is mainly produced industrially and exists naturally as a product of the decomposition of organic matter. It is also used as a refrigerant gas and in the production of plastics, textiles, dyes, explosives, and other chemicals. The emissions of NH₃ into the Earth's atmosphere have been increasing over the last few decades. The change in NH₃ concentration has essential implications for air quality and the ecosystem. To ascertain the significant influence of NH3 on many atmospheric reactions, several researchers have investigated the role of NH₃ on many important atmospheric reactions

(Jonas et al., 2013; Mallick et al., 2018; Ali, 2019; Kumar et al., 2020; Zhang et al., 2020; Ali et al., 2021). To this end, it is essential to clearly understand the reaction between *CH₂NH₂ and O₂ in the presence of NH₃, given its similar catalytic efficiency to water.

Water vapor is an environmentally significant constituent of the Earth's atmosphere. Numerous investigations have been carried out to determine the catalytic role of a single H₂O molecule in many atmospheric and combustion reaction systems (Vöhringer-Martinez et al., 2007; Iuga et al., 2010; Buszek et al., 2012; Iuga et al., 2011; Thomsen et al., 2012; Zhang et al., 2013; Zhang et al., 2014; Jara-Toro et al., 2017; Ali et al., 2018; Inaba, 2018; Ali et al., 2019; Ali, 2020; Wu et al., 2020; Ali and Balaganesh, 2022; Ali et al., 2022; Dash and Ali, 2022). These studies reveal that water-catalyzed reactions are energetically more favorable than other catalyzed reactions due to the formation of many hydrogen-bonded intermediates and transition states. However, water does not enhance the reaction's rate coefficients under tropospheric conditions due to its high concentration and lower entropic contribution compared to a free reaction (Ali et al., 2019; Ali, 2020; Ali and Balaganesh, 2022; Dash and Ali, 2022).

In this paper, we have investigated the rate coefficients for the effect of NH₃ and H₂O molecules on the important atmospheric and combustion prototype reactions, i.e., CH2NH2 + O2, for the first time. Using the RRKM/ME simulation, the temperature- and pressure-dependent rate coefficients were calculated between 200 K and 400 K and pressure ranges of 0.0001-1000 atm. The role of enthalpy and entropy contributions on hydrogen-bonded species on the effect of ammonia and water on the CH2NH2 + O2 reaction has been discussed to understand the chemical kinetic behavior of these complexes. In these situations, we have been inspired to model a gas-phase ternary reaction system, CH2NH2 $\cdots O_2 \cdots X$ (X = NH₃, H₂O), where H₂O and NH₃ can act as catalysts (vide Infra). To assess the accuracy of the data provided in this work, we have compared the energies and re-calculated rate coefficients and compared them with the available literature data for CH2NH2 + O₂ and its isoelectronic similar reaction, i.e., CH₂OH + O₂ (Dash and Ali, 2022). We hope that this study will strengthen the chemical kinetic database for global modeling and provide a thorough understanding for further study on analogous reaction systems.

2 Theoretical and computational methodology

2.1 Quantum chemical calculations

All the electronic structure calculations were carried out with the Gaussian 09 suite of programs (Frisch, 2013). The stationary points on potential energy surfaces (PESs) for ${}^{\bullet}CH_2NH_2 + O_2$, ${}^{\bullet}CH_2NH_2 + O_2$ (+NH₃), and ${}^{\bullet}CH_2NH_2 + O_2$ (+H₂O) reactions were computed using the hybrid-density functional method, i.e., M06-2X (Zhao and Truhlar, 2008) with the Pople 6-311++G (3df, 3pd) basis set (Frisch et al., 1984) and tabulated in Supplementary Table S1. The M06-2X is a frequently used preeminent functional to investigate the noncovalent interactions of transition states, intermediates, and post-intermediates for investigating chemical systems that encounter hydrogen bonding. To add corrections from the van der Waals interaction on M06-2X (Zhao and Truhlar, 2008), the Grimme

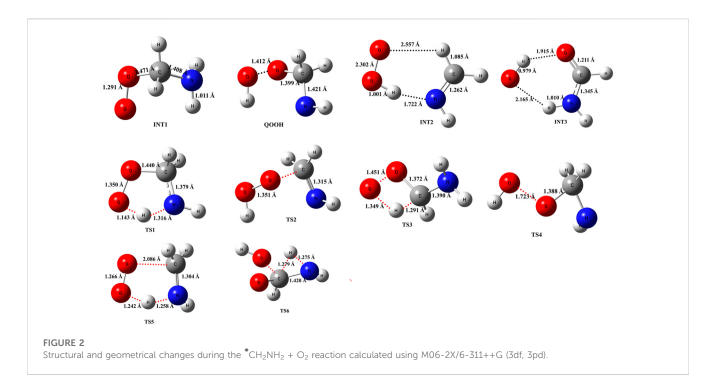
empirical dispersion "GD3" was used (Grimme et al., 2010). Normal modes of the vibrational frequency for each optimized species were carried out to obtain the zero-point energy (ZPE) and to calculate the rotational-vibrational partition functions. The transition state (TS) shows a single imaginary frequency, whereas reactants, intermediates, and products all show positive vibrational frequencies (see Supplementary Table S2). Intrinsic reaction coordinate (IRC) calculations (Fukui, 1981) were performed to confirm the identity of intermediates and post-intermediates for each TS. The IRC calculation was performed in both directions with the maxpoints=50 and the step size set to 3. The internal degrees of freedom of all species involved in the reaction were treated as harmonic oscillators and rigid rotor approximations, as suggested in previous studies for similar reaction systems (Ali et al., 2018; Dash and Ali, 2022). To improve the accuracy of energy, the single-point energy calculations were carried out at CCSD(T)/6-311++G (3df, 3dp)//M06-2X/6-311++G (3df, 3dp)+GD3 (Frisch et al., 1984; Raghavachari et al., 1989; Zhao and Truhlar, 2008). The result provides values that are accurate enough up to ~1 kcal/mol, as validated in our previous studies (Ali et al., 2019; Ali, 2020; Ali and Balaganesh, 2022; Dash and Ali, 2022). To check the qualitative contribution of the single-reference wave function, we have carried out the T1 diagnostic calculation at CCSD(T)/6-311++G (3df, 3pd). The calculated T1 diagnostic was found to be \leq 0.03, which is an acceptable range for a single reference wave function. To understand the spin contamination for each species, the spin expectation value <S2> was calculated and found to be in the range of ~0.75-0.77, which indicates that spin contamination was negligible.

2.2 State-of-the-art kinetics calculations

All the kinetics calculations were carried out using a software tool in the MultiWell suite of the program (Barker, 2009; Barker, 2011; Barker, 2023). The "me" codes in MultiWell programs calculate the unimolecular rate coefficients k(E) based on the RRKM/master equation as follows (Forst, 2003):

$$k(E) = \left[\frac{m^{\neq}}{m} \frac{\sigma_{ext}}{\sigma_{ext}^{\neq}}\right] \frac{g_e^{\neq}}{g_e} \frac{1}{h} \frac{G^{\neq}(E - E_{0,0})}{\rho(E)}.$$
 (1)

To avoid repetition from the previous studies, the details of each term of the equation are given in Supplementary Material S1. To calculate temperature- and pressure-dependent rate coefficients and branching fractions, N2 bath gases were used with an approximate value of the energy transfer process $\langle \Delta E \rangle$ $down = 200 \times (T/300)^{0.85} \text{ cm}^{-1}$ (Goldsmith et al., 2012). The Lennard-Jones parameters for collider gases (N_2) εk_B , $\sigma(N_2)$ = 3.74 Å, and $\varepsilon/k_B(N_2) = 82$ K were obtained from Hippler et al. (1983). The Lennard-Jones parameters of NH₂CH₂O₂ and NH₂CH₃O₂ were approximated based on Rissanen et al. (2014). The double arrays used in me simulations consisted of 1500 array elements with 10 cm⁻¹ energy grains using a quasi-continuum regime, which is evaluated up to 85,000 cm⁻¹. At each pressure and temperature value, ME simulations were carried out using the chemical activation energy distribution, which is appropriate for association reactions. The RRKM/ME simulations consisted of 10⁵ stochastic trials, each with a simulated time duration corresponding to an average of 100 collisions.



The pressure-dependent total rate coefficients $k^{bimol}(T, M)$ for ${}^{\bullet}\text{CH}_2\text{NH}_2 + \text{O}_2$ were calculated using (Ali, 2020; Dash and Ali, 2022)

$$k^{bimol}(T, M) = \Gamma K_{eq} \times k_{\infty}^{uni} (1 - f_{CH_2NH_2 + O_2}),$$
 (2)

where (Γ) is the quantum mechanical tunneling correction to the microcanonical rate coefficients k(E). Γ was implemented in the MultiWell master equation code, which is based on the 1-D Eckart asymmetric barrier. The k(E) calculated using the modified sums of states of the transition state reflect the tunneling effects. Tunneling was used to initialize the chemical activation distribution if both the "CHEMACT" and "TUN" keywords were selected. The $f_{CH_2NH_2+O_2}$ is the branching fraction (f) of the reaction going back to the reactants, and k_{∞}^{uni} is a high-pressure limit rate coefficient. The fall-off behavior of rate coefficients from (pressure = 1000 bar, P $\rightarrow\infty$) toward the low-pressure limit (p=0.0001 bar, P $\rightarrow0$) was considered

For the barrierless reactions, i.e., ${}^{\bullet}\text{CH}_2\text{NH}_2 + O_2 \rightarrow \text{CH}_2\text{NH}_2\text{OO},$ ${}^{\bullet}\text{CH}_2\text{NH}_2 \dots \text{H}_2\text{O} + O_2 \rightarrow \text{CH}_2\text{NH}_2\text{OO} \dots \text{H}_2\text{O}, {}^{\bullet}\text{CH}_2\text{NH}_2 \dots \text{NH}_3 + O_2 \rightarrow {}^{\bullet}\text{CH}_2\text{NH}_2\text{OO} \dots \text{NH}_3,$ the inverse Laplace transform (ILT) method was used. Since the rate coefficients for association reactions are usually weak and dependent on temperature, the activation energy for the recombination reaction was assumed to be equal to 0. As suggested in many similar reactions (Rissanen et al., 2014), this approach is good, and Arrhenius's activation energy can be equal to the reaction critical energy (E₀). MultiWell input for ILT calls for only two parameters E₀ and A-factor. In this work, we use statistical rate theories, which do not account for non-statistical effects, such as slow intramolecular vibrational energy redistribution (IVR), as suggested by Ali et al. (2023).

The equilibrium constant (K_{eq}) for the formation of ${}^{\bullet}CH_2NH_2 + O_2 \rightarrow CH_2NH_2OO^{\bullet}$, ${}^{\bullet}CH_2NH_2 \dots H_2O \rightarrow CH_2NH_2OO \dots H_2O$, ${}^{\bullet}CH_2NH_2 \dots NH_3 \rightarrow {}^{\bullet}CH_2NH_2OO \dots NH_3, {}^{\bullet}CH_2NH_2 \dots H_2O + O_2 \rightarrow CH_2NH_2OO \dots H_2O$ and ${}^{\bullet}CH_2NH_2 \dots H_2O$

 $NH_3+O_2 \rightarrow {}^{\bullet}CH_2NH_2OO \dots NH_3$ was calculated using the "THERMO" code as given: (Barker, 2009; Barker, 2011; Barker, 2023)

$$K_{eq} = \frac{Q_{INT}}{Q_R} exp\left(-\frac{E_{INT} - E_R}{k_B T}\right). \tag{3}$$

The equilibrium constants (K_{eq}) for the formation of two-body and three-body complexes calculated by Eq. 3 are tabulated in Supplementary Tables S3, S4. The Q_{INT} and Q_R are total partition functions of the intermediates and reactants, respectively; $E_{INT} - E_R$ is the zero-point corrected energy difference between intermediates and reactants. The calculated rate coefficients in the high-pressure limit (k_{∞}) were fitted to the modified Arrhenius expression $k_{\infty}(T) = A \times T^n \times \exp\left(\frac{-E_n}{RT}\right)$ in the temperature range of 200 K–400 K.

3 Results and discussion

3.1 Geometries and Energies

3.1.1 Reaction channels for ${}^{\circ}CH_2NH_2 + O_2$

The optimized structures of intermediates and transition states are shown in Figure 2. The zero-point corrected PES for the ${}^{\bullet}\text{CH}_2\text{NH}_2 + \text{O}_2$ reaction is depicted in Figure 3, and enthalpies values are given in Table 1. In the current reaction system, the O_2 molecule attacks the radical carbon atom, which leads to the formation of the intermediate ${}^{\bullet}\text{OO-CH}_2\text{NH}_2$ (INT1). Several conformational isomers of INT1 were observed, and for simplicity, we have considered the lowest energy conformer in our calculation. The calculated stabilization energy for INT1 is -31.7 kcal mol $^{-1}$, which is in very good agreement with the reported values by Rissanen et al. (2014) and Zhang et al. (2020) This value is also in very good agreement with its isoelectronic reactions, i.e., the $\text{O}_2 + {}^{\bullet}\text{CH}_2\text{OH}$ value (-31.9 kcal/mol) (Dash and Ali, 2022). The

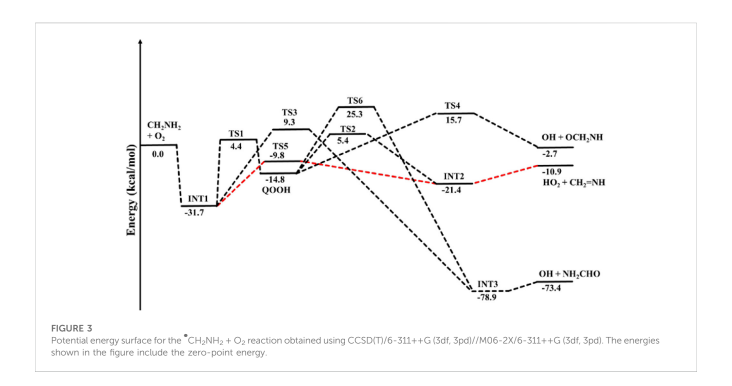


TABLE 1 Comparison of enthalpies (in kcal mol⁻¹) of each species for the *CH₂NH₂ + O₂ reaction with those found in previous studies and its isoelectronic analogs.

$^{\bullet}$ CH ₂ NH ₂ + O ₂ →	This Work	Previous works. Rissanen et al. (2014); Glarborg et al. (2020)	•CH ₂ OH + O ₂ →	Previous Work (Dash and Ali, 2022)	ΔS _{r,298K}
H ₂ NCH ₂ OO (INT1)	-31.7	-33.3 ^a ,-32.5 ^b	HOCH ₂ OO (Int-1)	-31.9	-37.3
H···NHCH ₂ OO (TS1)	4.4	2.5ª	H···OCH ₂ OO (TS-1)	-7.4	-39.6
HO ₂ ···CH ₂ NH (TS2)	5.4	2.7ª	HO ₂ ···CH ₂ O (TS-2)	2.8	-36.2
H…CHNH ₂ OO (TS3)	9.3	6.2ª	HO···HCOOH (TS-3)	8.6	-38.0
HO···OCH ₂ NH (TS4)	15.7	12.4ª	HO···OCH ₂ O (TS-4)	24.0	-35.3
H···NHCH ₂ ···OO (TS5)	-9.8	-10.2 ^a ,-12.0 ^b	H···O ₂ ··CH ₂ O (TS-5)	-18.5	-38.9
HNHCHOOH (TS6)	25.3	21.8ª	-	-	-35.9
HO···OCH ₂ NH (QOOH)	-14.8	-16.1ª	HO···OCH ₂ O	-14.2	-37.6
OOH···NHCH ₂ (INT2)	-21.4	-22.8 ^a ,22.0 ^b	OOH···OCH ₂ (Int-2)	-25.6	-29.2
HO···NH ₂ CHO (INT3)	-78.9	-77.2 ^a	-	-	-29.3
CH ₂ NH + HO ₂	-10.9	-12.2 ^a ,-10.6 ^b	CH ₂ O+ HO ₂	-17.9	1.8
NH ₂ CHO + OH	-73.4	-73.1ª			-2.0
OCH ₂ NH + OH	-2.7	-2.9ª	OCH ₂ O+ OH	-15.0	-5.5

^aRissanen et al. (2014).

^bGlarborg et al. (2020).

unpaired electron in INT1 resides at the terminal O-atom, which can be decomposed differently. The lowest energy channel is an isomerization process where the terminal O-atom attacks the H-atoms of the NH_2 group via five- and six-membered cyclic transition states (TS5 and TS1),

leading to hydrogen-bonded five- and six-membered cyclic complexes, i.e., INT2 and QOOH, respectively. The calculated barrier heights for TS5 and TS1 are 21.9 and 36.1 kcal mol⁻¹, respectively, with respect to INT1, indicating that the isomerization reaction going through TS5,

leading to the formation of INT2, is energetically more favorable than that going through TS1 to QOOH. Supplementary Figure S1 provides an IRC scan that confirms the connectivity of TS5 with INT2 and CH2NH + HO₂ at the M06-2X/6-311++G (3df, 3pd). It is noted that the influence of the formation of hydrogen-bonded cyclic complexes may change the energetics and kinetics of the reaction system. In INT2, two strong hydrogen bonds are formed between the H-atom of the HO2 and the N-atom of the CH₂NH (1.72 Å) and O-atom of the HO₂ radical and the H-atom of the CH₂NH (2.55 Å), which leads to the formation of a stable six-membered ring planar cyclic structure. In QOOH, a five-membered ring cyclic structure with one hydrogen bond is formed between the terminal O and H-atoms (2.47 Å). Therefore, INT2 is energetically 6.6 kcal mol⁻¹ more stable than QOOH. QOOH can further dissociate via TS2 and TS4 to form HO2 + CH2NH and OH + OCH2NH, respectively. Because the barrier height for the formation of QOOH is very high, the formation of CH2NH and OCH2NH via QOOH may be negligible under tropospheric conditions. The terminal O-atom of INT1 can also attack the H-atom of a nearby C-atom, leading to the formation of a hydrogen-bonded six-membered cyclic intermediate, INT3, via a four-membered ring transition state (TS3) and subsequently dissociating to form OH + NH₂CHO. In INT3, two strong hydrogen bonds are formed between the H-atom of the OH radical and the O-atom of the NH₂CHO (1.91 Å) and the O-atom of the OH radical and the H-atom of the NH₂CHO (2.16 Å). INT3 is energetically the most stable structure in the PES, with a stabilization energy of -78.9 kcal mol⁻¹ from the reactants. The barrier height of this reaction channel is 41 kcal mol⁻¹, which is 20 kcal mol⁻¹ higher than that of the TS5 and may not contribute to the overall reaction kinetics (vide infra).

The other reaction channel, such as the conversion from QOOH to INT3 via H-atom shift (C to N) through a three-membered transition state, TS6 (41 kcal mol⁻¹), is expected to have a negligible impact on the total rate coefficient due to its high energy barriers.

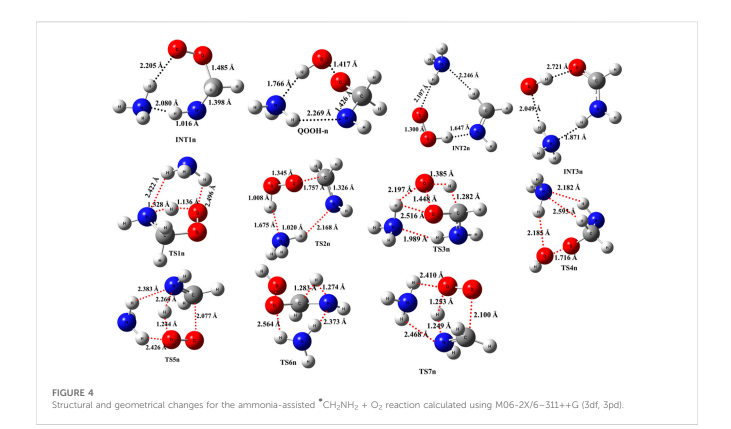
The enthalpies of reaction (ΔH_{rxn} (0 K) for ${}^{\bullet}CH_2NH_2 + O_2 \rightarrow$ CH₂NH + *HO₂ (-10.9 kcal mol⁻¹) are in very good agreement with those found in the most accurate active thermochemical database (ATcT) (-11.42 kcal mol⁻¹) (Ruscic et al., 2004; Ruscic and Bross, 2020) and in good agreement with the theoretically calculated value in Rissanen et al. (2014) (-12.2 kcal mol⁻¹). The computed PES for the ^oCH₂NH₂ + O₂ reaction is also consistent with its isoelectronic analogous reaction system, i.e., CH2OH + O2 reported by Dash and Ali (2022) using CCSD(T)//ωB97XD/6-311++G (3df, 3pd) level of theory. The enthalpy values obtained in their calculations are also given in Table 1. The reaction energies for $CH_3NH^{\bullet} + O_2 \rightarrow CH_2NH +$ HO₂ (-17.3 kcal/mol) are also calculated and found to be in very good agreement with that in the ATcT (-17.9 kcal/mol) (Ruscic et al., 2004; Ruscic and Bross, 2020). The energies obtained for most of the structures of the CH₂OH + O₂ system (Ali, 2020) along the reaction paths are very close to those of the current system, indicating the reliability of the data presented here. However, the barrier heights for the isomerization pathways Int-1→TS-1 (-7.4 kcal/mol)→QOOH and Int-1 →TS-5 (-18.5 kcal/mol) Int-2 in the CH₂OH + O₂ system (Dash and Ali, 2022) are quite low and more stable compared to those in the same pathways in the current system with respect to the reactant's energy (Table 1). Moreover, the barrier height for the reaction proceeding from HO···OCH₂O to OCH₂O + OH (Dash and Ali, 2022) is 24 kcal/mol (TS-4), and the stabilization energy for the products is -15 kcal/mol, whereas in the current system, the corresponding energies are 8.3 and 12.3 kcal/mol less stable than the former ones, respectively. These differences in barrier energies can affect the overall rate coefficients between the two systems.

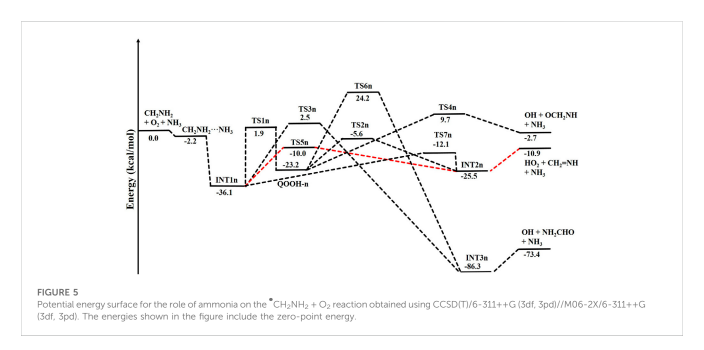
3.1.2 Role of the ammonia molecule on ${}^{\bullet}CH_2NH_2 + O_2$

When a single ammonia molecule is introduced in ${}^{\bullet}\text{CH}_2\text{NH}_2 + O_2$, the simultaneous collision between ${}^{\bullet}\text{CH}_2\text{NH}_2$, O_2 , and NH₃ is very unlikely to occur; therefore, the probability of a trimolecular reaction is very small under real conditions. Hence, the first step is the formation of a CH₂NH₂...NH₃ complex, followed by collision with O₂. The CH₂NH₂...NH₃ complex (-2.2 kcal/mol) is assumed to be more important than CH₂NH₂...O₂ and NH₃...O₂ due to its lower binding energy (<1 kcal/mol). As discussed in our previous work, we have also used a similar approach for ammonia-assisted reactions (Ali et al., 2019; Ali, 2020; Ali and Balaganesh, 2022; Dash and Ali, 2022). The geometrical changes in ammonia-assisted intermediates and transition states are shown in Figure 4, and the zero-point corrected PES for the ammonia-assisted ${}^{\bullet}\text{CH}_2\text{NH}_2 + O_2$ reaction is given in Figure 5. The energy of all the stationary points, i.e., reactants, intermediates (INTs), and transition states, is tabulated in Table 2.

Figure 5 shows that the effect of the ammonia reaction proceeds via similar reaction pathways as a free reaction. For simplicity, only the most stable structures are shown in the PES. As shown in Figure 5, O₂ attacks the bimolecular complex CH₂NH₂···NH₃ to form a trimolecular hydrogen-bonded complex (INT1n) (Figure 4), whose stabilization energy is -36.1 kcal mol⁻¹. The resulting ammonia-assisted intermediate (INT1n) is 4.4 kcal mol⁻¹ more stable than the corresponding ammonia-free intermediate, i.e., INT1. This is due to the formation of strong hydrogen bonds between the terminal O-atom of H2NCH2OO and one of the H-atoms of NH₃ (2.20 Å) and the H-atom of H₂NCH₂OO (2.08 Å) with the N-atom of NH₃, whereas no such effect is observed in INT1. On the other hand, Table 2 shows that INT1 is entropically more favorable than INT1n with respect to reactants. This is due to the fact that the hydrogen-bonded complex decreases the entropy of the system. Similar to uncatalyzed reaction pathways, the terminal O-atom intra-molecularly attacks the H-atoms in the NH2 group in the presence of NH3, leading to the formation of INT2n and QOOH-n via five-membered cyclic transition states (TS5n/TS7n and TS1n, respectively). The difference in the barrier heights between two isomeric transition states, TS5n and TS7n, is 2.1 kcal mol⁻¹. TS5n seems to be more stable than TS7n because, in the case of TS5n, all three hydrogen atoms of ammonia face toward the molecular center, leading to the formation of a sixmembered ring hydrogen-bonded cyclic structure, whereas in the case of TS7n, hydrogen atoms of ammonia are away from the molecular center, leading to the formation of a similar sixmembered ring hydrogen-bonded cyclic structure. Entropy data also support that TS5n is more disordered than TS7n.

Supplementary Figure S2 provides an IRC scan at the same level that confirms that TS5n bridges the OCH₂C(O)OOH radical (INT2n) and CH₂NH + HO₂ +NH₃. The IRC scan confirms that the only stationary point between INT2h and the trimolecular products is that associated. The barrier height of TS1n is 2.5 kcal mol⁻¹ lower than that of the corresponding ammonia-free transition state TS1. This is due to the formation of two strong hydrogen bonds (H-atoms of ammonia with N and O atoms of the cyclic ring) in TS1n (1.93 and 2.14 Å). On





the other hand, the barrier height of TS5n is almost similar to that of TS5, although hydrogen bonds are present in TS5n. The differences in the barrier height can be explained by the formation of two adjacent cyclic ring structures (five- and six-membered), as aforementioned in TS5n, making it sterically hindered compared to only one ring structure in TS5. The stabilization energies of ammonia-assisted QOOH and INT2 are calculated to be -23.3 and -25.5 kcal mol⁻¹, respectively, which are 8.4 and 4.1 kcal mol⁻¹ lower than those of the corresponding free-ammonia structures. This can be explained

similarly by comparing the presence of hydrogen bonds in the respective structures. In QOOH-n, the formation of two strong hydrogen bonds between the H-atom of ammonia with N atom QOOH (2.26 Å) and N atom NH $_3$ and terminal H atom of QOOH (1.76 Å) leads to a seven-membered ring-like structure rather than only one hydrogen bond in the case of free QOOH. Similarly, the stability of INT2n can be explained by the formation of an eightmembered ring with three hydrogen bonds, H of HO $_2$ and N of CH $_2$ NH (1.64 Å), O of HO $_2$ and N of NH $_3$ (2.1 Å), and H of CH $_2$ NH

TABLE 2 Enthalpies (in kcal mol $^{-1}$) and entropies (in cal K $^{-1}$ mol $^{-1}$) due to the effect of NH $_3$ on each species involved for the $^{\circ}$ CH $_2$ NH $_2$ + O $_2$ reaction.

$^{\bullet}CH_{2}NH_{2} + O_{2} (+NH_{3}) \rightarrow$	ΔH _{rxn} (0 κ)	ΔS _{rxn} (298 κ)
CH ₂ NH ₂ ···NH ₃	-2.2	-28.2
H ₂ NCH ₂ OO···NH ₃ (INT1n)	-36.1	-66.3
H···NHCH ₂ OO···NH ₃ (TS1n)	1.9	-68.9
HO ₂ ···CH ₂ NH···NH ₃ (TS2n)	-5.6	-70.4
H···CHNH ₂ OO···NH ₃ (TS3n)	2.5	-70.9
HO···OCH ₂ NH···NH ₃ (TS4n)	9.7	-66.7
H···NHCH ₂ ···OO···NH ₃ (TS5n)	-10.0	-68.6
H···NHCHO···OH···NH ₃ (TS6n)	24.2	-66.5
H···NHCH ₂ ···OO···NH ₃ (TS7n)	-12.1	-69.6
HO···OCH ₂ NH···NH ₃ (QOOH-n)	-23.2	-68.8
OOH···NHCH ₂ ···NH ₃ (INT2n)	-25.5	-55.5
HO···NH ₂ CHO (INT3n)	-86.3	-61.2

and N of $\mathrm{NH_3}$ (2.24 Å) compared to a six-membered ring with two hydrogen bonds in the case of INT2. On the other hand, the structures of QOOH-n and INT2n are entropically less favorable compared to those of uncatalyzed QOOH and INT2n.

QOOH-n further dissociates to INT2n (via TS2n) and INT3n (via TS6n) and then subsequently forms $HO_2+CH_2NH + NH_3$ and OH +OCH₂NH + NH₃ via TS4n and OH + NH₂CHO + NH₃. INT2n and INT3n are eight-membered ring hydrogen-bonded structures, and their stabilization energies are 4.1 kcal mol⁻¹ and 7.4 of kcal mol⁻¹ lower than those of the corresponding uncatalyzed intermediates. Ammonia-assisted intermediates are more stable than ammonia-free ones because of the formation of an eight-membered ring structure with three strong hydrogen bonds between HO₂···NH₃ (2.1 Å), NH₃···CH₂NH (2.24 Å), and HO₂···CH₂NH (1.64 Å) in INT2n and OH···NH₃ (2.04 Å), $NH_3...NH_2CHO$ (1.87 Å), and $HO...NH_2CHO$ (1.72 Å) in INT3n. The barrier heights of TS2n (-5.6 kcal mol⁻¹), TS3n (2.5 kcal mol⁻¹), TS4n (9.7 kcal mol⁻¹), and TS6n (24.2 kcal mol⁻¹) were also consistently lower than those of the corresponding ammonia-free transition states due to similar hydrogen bonding interactions. Overall, the reaction in the presence of an ammonia-assisted intermediate is thermodynamically more favorable than the free reaction, and vice versa entropically.

3.1.3 Role of the water molecule on ${}^{\circ}CH_2NH_2 + O_2$

As previously discussed in the case of ammonia reactions, we have also employed a similar approach for water reactions. When a single H_2O molecule is added to ${}^{\bullet}CH_2NH_2 + O_2$, the first step is the formation of a $CH_2NH_2\cdots H_2O$ complex, followed by collision with O_2 . The $CH_2NH_2\cdots H_2O$ (-2.7 kcal/mol) is assumed to be more important than $CH_2NH_2\cdots O_2$ and $H_2O\cdots O_2$ (<1 kcal/mol) due to lower binding energy. The geometrical changes in water-assisted intermediates and transition states are shown in Figure 6 and the Cartesian coordinates of all the optimized geometries are given in Supplementary Table S1.

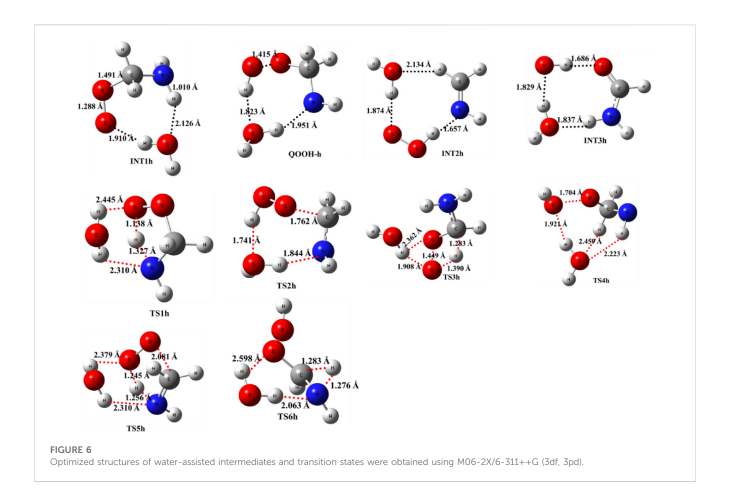
The zero-point-corrected PES for the water-assisted ${}^{\bullet}CH_2NH_2 + O_2$ reaction is given in Figure 7, and the energy of all the stationary points, i.e., reactants, INTs, and TSs, is tabulated in Table 3. As shown in

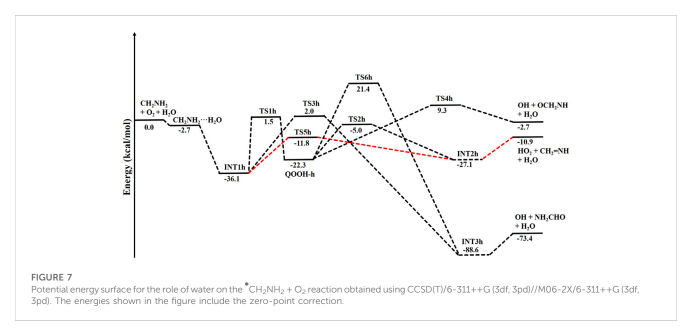
TABLE 3 Enthalpies (in kcal mol $^{-1}$) and entropies (in cal K $^{-1}$ mol $^{-1}$) due to the effect of H $_2$ O on each species involved for the $^{\circ}$ CH $_2$ NH $_2$ + O $_2$ reaction.

*CH ₂ NH ₂ + O ₂ (+H ₂ O) →	ΔH _{rxn} (0 κ)	ΔS _{rxn} (298 κ)
CH ₂ NH ₂ ···H ₂ O	-2.7	-27.9
H ₂ NCH ₂ OO··· H ₂ O (INT1h)	-36.1	-70.3
H···NHCH ₂ OO··· H ₂ O (TS1h)	1.5	-71.7
HO ₂ ···CH ₂ NH···NH ₃ (TS2h)	-5.0	-71.5
H···CHNH ₂ OO···NH ₃ (TS3h)	2.0	-72.0
HO···OCH ₂ NH···NH ₃ (TS4h)	9.3	-70.9
H···NHCH ₂ ···OO···NH ₃ (TS5h)	-11.8	-70.4
H···NHCHO···OH···NH ₃ (TS6h)	21.4	-66.2
HO···OCH ₂ NH···NH ₃ (QOOH-h)	-22.3	-71.4
OOH···NHCH ₂ ···NH ₃ (INT2h)	-27.1	-60.1
HO···NH ₂ CHO (INT3h)	-88.6	-64.1

Figure 6 and Figure 7, the O_2 molecule attacks the bimolecular complex ${}^{\circ}\text{CH}_2\text{NH}_2\text{---}\text{H}_2\text{O}$ to form a trimolecular hydrogen-bonded complex (INT1h) with a stabilization energy of -36.1 kcal mol^{-1} . The INT1h is 4.4 kcal mol^{-1} lower than that of the water-free intermediate (INT1) and identical to the energy of ammonia-assisted INT1n. The result indicates water- and ammonia-assisted reactions are energetically more favorable than free reactions.

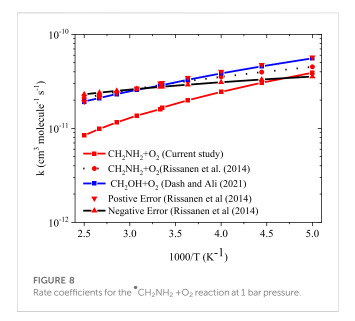
In INT1h, two strong hydrogen bonds are observed between the terminal O-atom of H2NCH2OO and one of the H-atoms of H2O (1.91 Å) and O-atom H₂O and terminal H-atom H₂NCH₂OO (2.12 Å) (see Figure 6). On the other hand, Table 3 and Table 1 show that INT1h is entropically least favored than INT1n and INT1. Similar to the free reaction, the terminal O-atom of INT1h intra-molecularly attacks the H-atoms in the NH₂ group, leading to the formation of cyclic structures, i.e., INT2h and QOOH-h, via five-membered cyclic transition states, i.e., TS5h and TS1h, respectively. The barrier heights of TS5h and TS1h are 2 kcal mol⁻¹ and 3 kcal mol⁻¹ lower than those of water-free transition states TS5 and TS1, respectively. In a similar manner, TS5h and TS1h are 1.8 kcal mol⁻¹ and 0.4 kcal mol⁻¹ lower than TS5n and TS1n, respectively. These differences in barrier heights indicate that water-assisted transition states are energetically more stable than ammonia-assisted and water-free species. The stabilization energies of water-assisted QOOH-h and INT2h are calculated to be -22.3 kcal mol⁻¹ and -27.1 kcal mol⁻¹, which are and 5.7 kcal mol⁻¹ lower than those of the 7.8 kcal mol⁻¹ corresponding free structures. This can be understood by comparing the presence of hydrogen bonds in the respective structures. Supplementary Figure S3 provides an IRC scan at the same level that confirms TS5h bridges the OCH2C(O)OOH radical (INT2h) and CH₂NH + H₂O + HO₂ products. The IRC scan confirms that the only stationary point between INT2h and the trimolecular products is that associated. In QOOH-h, the formation of two strong hydrogen bonds between the H-atom of water with the N atom of QOOH (1.95 Å) and the O atom of H₂O and the terminal H atom of QOOH (1.82 Å) led to the formation of a seven-membered ring-like structure. Similarly, the stability of INT2h can be found by the formation of an eight-membered cyclic ring with three hydrogen bonds, H of HO2 and

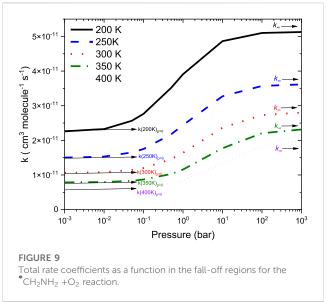




N of CH₂NH (1.65 Å), O of HO₂ and H of H₂O (1.87 Å), and H of CH₂NH and O of H₂O (2.13 Å), which are compared to that of a six-membered ring with two hydrogen bonds, as indicated in the case of INT2. Between water- and ammonia-catalyzed QOOH and INT2, QOOH-n is more stable than QOOH-h and INT2h is more stable than INT2n.

In general, the water-free pathways are entropically more favorable than water-free ones. The QOOH further dissociates to INT2h (via TS2h) and INT3h (via TS6h) and then subsequently forms $\rm HO_2 + CH_2NH + H_2O$, $\rm OH + OCH_2NH + H_2O$ via TS4h, and $\rm OH + NH_2CHO + H_2O$. In general, the water-assisted reaction channels are thermodynamically more favorable and entropically





less favorable than the free reaction. It is also clear from Table 3 that all other pathways are thermodynamically less important compared to $R + O_2 \rightarrow INT1h \rightarrow TS5h \rightarrow INT2h \rightarrow CH_2NH + HO_2 + H_2O$, whose barrier height is the lowest with respect to reactants. Therefore, the other reaction channels may have less contribution under tropospheric conditions.

3.2 Kinetics

3.2.1 Rate coefficients for the ${}^{\bullet}CH_2NH_2 + O_2$ reaction

To obtain the rate coefficients for ${}^{\bullet}CH_2NH_2 + O_2$ in temperatures between 200 K and 400 K and pressures from 0.000001 bar to 1000 bar, the RRKM/ME simulation has been used. The rate coefficients as a function of the temperature at 1 bar pressure are shown in Figure 8. The rate coefficients for the formation of O₂-CH2NH and CH2NH are observed to be pressure-dependent and negative temperature-dependent. This result is consistent with that of the previous reports by Rissanen et al. (2014) and a similar reaction system, i.e., CH2OH + O2 (Dash and Ali, 2022). The calculated rate coefficient at 300 K (1.6 × 10⁻¹¹ cm³ molecule⁻¹ s⁻¹) is a factor of ~2 lower than the experimentally measured ones $(3.2 \times 10^{-11} \, \text{cm}^3)$ molecule⁻¹ s⁻¹) (Rissanen et al., 2014). The calculated value is in good agreement with the lower temperature range <250 K when the positive error is considered. In light of the expected errors in computed thermochemistry, which may be up to 1 kcal/mol, we believe that this level of accuracy is sufficient for these purposes. A recent study by Ali et al. (2023) indicates that the barrier heights of similar reactions are very sensitive to quantum chemical calculations. The computed rate coefficients are compared with previously reported values and also with its isoelectronic analogous reaction system, i.e., CH₂OH + O₂ (see Figure 8) (Dash and Ali, 2022). The calculated values at 300 K are a factor of 2 lower than the theoretically calculated values (Dash and Ali, 2022). The presence of the N atom in CH₂NH and the O atom ^oCH₂OH can explain this, leading to the development of various chemical kinetics conclusions.

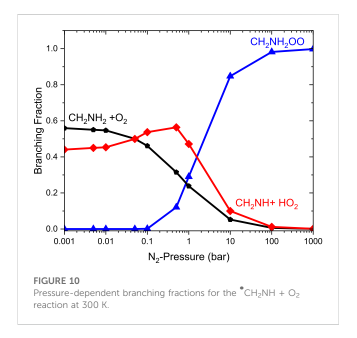


Figure 9 reports the rate coefficients in the fall-off regions for the ${}^{\circ}\text{CH}_2\text{NH}_2 + \text{O}_2$ reaction at different temperatures. The rate coefficients increase as pressure increases, and the HPL condition is observed at ~100 bar. As shown in Figure 9, the largest difference between the two limits occurs at about 250 K, which reaches a factor of 2. However, at 200 K, the difference between the two regimes is about a factor of 2. To provide more detailed insights, the relative branching fractions of these channels were determined at 200 K, 300 K, and 400 K and shown in Supplementary Figure S4. For simplicity, the branching fraction at different pressures and at 300 K is shown in Figure 10. At all temperatures and pressures, the branching fractions of QOOH are almost negligible; therefore, they are not shown in Figure 10. The branching fraction for the formation CH₂NH/HO₂ contributes 40% at 300 K and increases as temperature increases to 400 K. The formation of CH₂NH decreases

TABLE 4 Calculated rate coefficients for the $^{\circ}$ CH₂NH₂ + O₂, $^{\circ}$ CH₂NH₂ +O₂ (+NH₃), and $^{\circ}$ CH₂NH₂ +O₂ (+H₂O) in the temperature range of 200 K–400 K at 1 bar pressure.

Temperature	*CH ₂ NH ₂ + O ₂	Exp. value Rissanen et al. (2014)	³² CH ₂ OH + O ₂	*CH ₂ NH ₂ +O ₂ (+NH ₃)	*CH ₂ NH ₂ +O ₂ (+H ₂ O)
200	3.9×10^{-11}	1.9×10^{-11}	5.6×10^{-11}	1.9×10^{-22}	9.8×10^{-20}
225	3.1×10^{-11}	2.2×10^{-11}	4.6×10^{-11}	8.6×10^{-23}	7.8×10^{-19}
250	2.4×10^{-11}	2.4×10^{-11}	3.8×10^{-11}	4.8×10^{-23}	3.7×10^{-18}
275	2.0×10^{-11}	2.7×10^{-11}	3.3×10^{-11}	3.0×10^{-23}	1.3×10^{-17}
298	1.7×10^{-11}	2.9 × 10 ⁻¹¹	2.8×10^{-11}	1.9×10^{-23}	3.3×10^{-17}
300	1.6×10^{-11}	2.8×10^{-11}	2.85×10^{-11}	1.9×10^{-23}	3.3×10^{-17}
325	1.4×10^{-11}	3.1×10^{-11}	2.6×10^{-11}	1.4×10^{-23}	7.4×10^{-17}
350	1.2×10^{-11}	3.4×10^{-11}	2.3×10^{-11}	1.1×10^{-23}	1.4×10^{-16}
375	0.9×10^{-11}	3.6×10^{-11}	2.1×10^{-11}	8.7×10^{-24}	2.3×10^{-16}
400	0.8×10^{-11}	3.8×10^{-11}	1.9×10^{-11}	7.4×10^{-24}	4.0×10^{-16}
k = AT ⁿ	A = 0.02		$A = 1.0 \times 10^{-09}$	$A = 4.8 \times 10^{-31}$	$A = 3.5 \times 10^5$
	n = -3.4		n = -0.75	n = 2.0	n = -6.0
exp (-B/T)	B = 323		B = -222	B = -1858	B = 4958

to 0% as pressure increases to 100 bar. At the same time, the formation of $\rm CH_2NH_2OO$ increases as pressure increases to 100 bar (100%). The plot shows that backward reaction to regenerate reactants is prominent at <0.01 bar. These results are also consistent with the previously reported branching ratio by Rissanen et al. (2014). At all the pressure rates, it is more advantageous to lose $\rm HO_2$ via the formation of $\rm CH_2NH$ rather than through OH loss via formamide formation due to an energetically favorable pathway (see Figure 10). As temperature increases from 300 K to 400 K, the product branching ratio increases (see Supplementary Figure S4). The result is due to the fact that the relevant stationary points on the respective potential energy surfaces have a dominant entropy factor over the enthalpy factor.

3.2.2 Rate coefficients for *CH₂NH₂+O₂ (+NH₃)

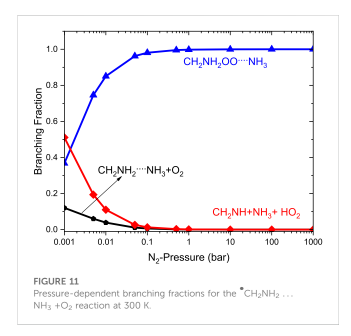
As discussed in the previous section, only the entry channel ${}^{\bullet}\text{CH}_2\text{NH} \dots \text{NH}_3 + \text{O}_2$ is considered for the rate coefficient calculations.

$$\begin{array}{c} CH_2NH_2+O_2+NH_3 \overset{k_{1n}}{\underset{k_{-1n}}{\rightleftarrows}} CH_2NH_2---NH_3+O_2 \\ \overset{k_{2n}}{\underset{k_{-2n}}{\rightleftarrows}} O_2-CH_2NH_2---NH_3 \overset{k_{uni-n}}{\longrightarrow} CH_2NH+HO_2+NH_3 \end{array}$$

The equation to calculate the effective pressure-dependent rate coefficients $k_{eff}^{bimol}(T,M)$ is as follows:

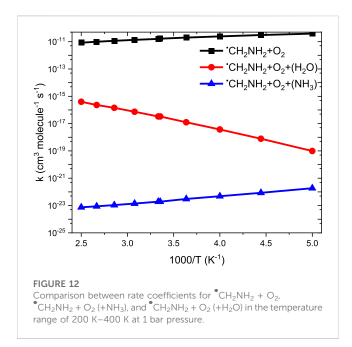
$$k_{eff}^{bimol}(T, M) = K_{eq-1n} \times [NH_3] \times K_{eq-2n} \times k_{\infty}^{uni} \left[1 - f_{(\bullet CH_2 NH_2 ... NH_3 + O_2)}\right], \tag{4}$$

where $K_{eq-1h} = \frac{k_{1n}}{k_{-1n}}$ and and $K_{eq-2n} = \frac{k_{2n}}{k_{-2n}}$ are equilibrium constants of each pathway involved in a reaction, [NH₃] is the concentration, and f is the branching fraction for the reaction proceeding to the reactant. The ammonia concentration used at 10 ppbv is based on the observations from previous studies (Ali et al., 2021). The rate coefficients for the ammonia-assisted



reaction are almost similar to those of ${}^{\bullet}\text{CH}_2\text{NH}_2 + \text{O}_2$ in the temperature range of 200 K–400 K (see Table 4). In fact, some lower values were obtained at higher temperatures. The catalytic behavior does not take place if step 0 is not included in the reaction mechanism. The result could be due to the lower entropy change in the reaction. The total effective rate coefficient for ${}^{\bullet}\text{CH}_2\text{NH}_2 + \text{O}_2 \ (2.7 \times 10^{-21} \ \text{cm}^3 \ \text{molecule}^{-1} \ \text{s}^{-1} \ \text{at } 300 \ \text{K})$ is $\sim 12 \ \text{orders}$ of magnitude lower than that for the ${}^{\bullet}\text{CH}_2\text{NH} + \text{O}_2 \ \text{reaction} \ (8.8 \times 10^{-13} \ \text{cm}^3 \ \text{molecule}^{-1} \ \text{s}^{-1})$. This result is due to the fact that the ammonia-assisted pathway depends on the ammonia concentration (see Table 4).

The relative branching fractions of these channels determined at 200 K, 300 K, and 400 K are shown in Supplementary Figure S5. For



simplicity, the branching fraction at 300 K is shown in Figure 11. The branching fraction for the formation of CH_2NH/HO_2 contributes 50% at 300 K, and almost the same temperature increases to 400 K (see Supplementary Figure S5). At the same time, the formation of $CH_2NH_2OO \dots NH_3$ increases as the pressure increases to 100 bar (100%). The plot shows the backreaction that regenerates ${}^{\bullet}CH_2NH_2\dots NH_3 + O_2$. When the results are compared with those of the free reaction, it is easier to lose HO_2 via the formation of CH_2NH than through OH loss via formamide formation. We can say that the effect of ammonia has a negligible impact on the product branching ratios, and the results are almost similar to those of a free reaction, except at very low pressure (Supplementary Figure S5).

3.2.3 Rate constant for ${}^{\bullet}CH_2NH_2 + O_2 (+H_2O)$

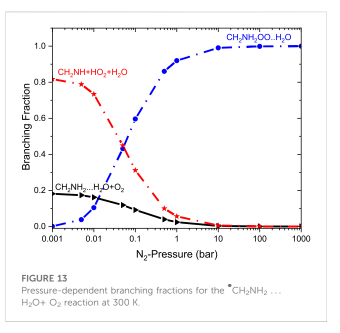
The scheme for the formation of INT1h, CH_2NH , and HO_2 from ${}^{\bullet}CH_2NH_2 + O_2$ reactions with the effect of a water can be written as follows.

$$\begin{array}{c} CH_2NH_2+O_2+H_2O \underset{k_{-1h}}{\overset{k_{1h}}{\longleftrightarrow}} CH_2NH_2---H_2O+O_2 \\ \overset{k_{2h}}{\longleftrightarrow} O_2-CH_2NH_2---H_2O \xrightarrow{k_{uni-h}} CH_2NH+HO_2+H_2O \end{array}$$

The equation to calculate the effective pressure-dependent rate coefficients $k_{eff}^{bimol}(\exp(-B/T))$ is as follows:

$$k_{eff}^{bimol}(T, M) = K_{eq-1h} \times [H_2O] \times K_{eq-2h} \times k_{co}^{uni} \left[1 - f_{(\bullet CH_2NH_2...H_2O+O_2)}\right], \tag{5}$$

where $K_{eq(1)h} = \frac{k_{1h}}{k_{-1h}}$ and $K_{eq-2h} = \frac{k_{2h}}{k_{-2h}}$ are the equilibrium constants of each reaction pathway involved in equation (iii), $[H_2O]$ is the concentration, and f is the branching fraction for the reaction proceeding to the reactants. The $[H_2O]$ is calculated using a typical humidity concentration, as discussed in the previous paper (Dash and Ali, 2022). The rate coefficients were also calculated using different water concentrations, as shown in Supplementary Figure S6. The effect of relative

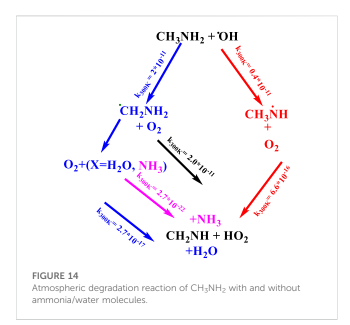


humidity from 20% to 100% on calculated rate coefficients is a factor of 10 difference. The effective rate coefficient calculated based on Eq. 4 (2.04×10^{-17} cm³ molecule $^{-1}$ s $^{-1}$ at 298 K) is ~six to seven orders of magnitude lower than that of the water-free $^{\bullet}$ CH₂NH₂ + O₂ reaction ($^{\sim}$ 2.2 × 10^{-11} cm³ molecule $^{-1}$ s $^{-1}$ at 298 K). This is due to the fact that the water-assisted pathway depends parametrically on water concentration and entropy reduces the rate coefficients. Our results are also consistent with the previously reported values for similar reaction systems (Ali et al., 2022; Dash and Ali, 2022).

Our calculation shows that the total effective rate coefficients for systems ${}^{\bullet}CH_2NH_2 + O_2$ (+NH₃) (~10⁻¹¹ order) and ${}^{\bullet}CH_2NH_2 + O_2$ (+H₂O) (6 order) are smaller than that of the free reaction (see Figure 12).

It is clear that the geometries of INT and TSs are different in [•]CH₂NH₂ + O₂ (+NH₃) reaction systems compared to their isoelectronic analogous CH2NH2 + O2 (+H2O) reactions, resulting in different computed enthalpies and rate coefficients. Because of this, the kinetics of CH₂NH₂ + O₂ (+NH₃) is quite different from those of $^{\bullet}$ CH₂NH₂ + O₂ (+H₂O) reaction systems. In the case of free reactions and ammonia, the rate coefficients exhibit negative temperature dependence, whereas in the case of water, positive temperature dependence was observed. This may be due to the fact that water concentration is highly dependent on temperature, and ammonia concentration is nearly independent of temperature. The branching fractions for the formation of *OOCH2NH2 and *CH2NH and the reaction going back to CH2NH2 + O2 with the effect of a single water molecule at 300 K and pressure range 0.001 bar-1000 bar are shown in Figure 13, and a comparison of branching fractions for CH2NH2 + O_2 , ${}^{\bullet}CH_2NH_2 + O_2$ (+NH₃), and ${}^{\bullet}CH_2NH_2 + O_2$ (+H2O) at 200 K, 300 K, and 400 K is shown in Supplementary Figure S7.

As previously discussed in water- and ammonia-free reactions, the product branching ratios for the formation of $CH_2NH + HO_2$ decrease, and the reaction goes back to the reactants, i.e., ${}^{\bullet}CH_2NH_2 + O_2$, when the pressure increases from 0.1 bar. When a water molecule is added to the reaction, the product branching ratio changes significantly (~80%), and a single water reaction favors



the formation of $\mathrm{CH_2NH} + \mathrm{HO_2}$ at a temperature of <300 K; however, the effect of ammonia favors only ~10%. Despite the slower water reaction, our ME calculations indicate that a favorable $\mathrm{CH_2NH} + \mathrm{HO_2}$ formation is observed under tropospheric conditions.

3.3 Atmospheric fate of methylamine and methanimine

The atmospheric degradation of CH_3NH_2 with and without ammonia and water molecules is shown in Figure 14. The atmospheric lifetime (τ) because of its interaction with OH and species ${}^{\bullet}CH_2NH_2$ with O_2 radicals is calculated as follows:

$$\tau = \frac{1}{k_{[X=OH,O2]} \times [X]}.$$
 (6)

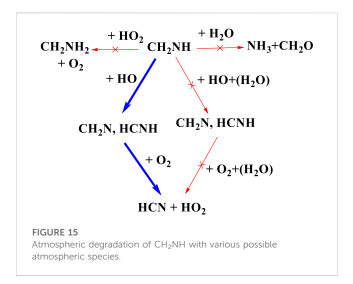
The average OH radical concentration at tropospheric conditions 225 K and <1 bar is $\sim 1 \times 10^6$ cm³ molecule⁻¹ s⁻¹, and a concentration of $[O_2] = 1 \times 10^{16}$ molecules cm⁻³ was used, which is based on a previous study (Dash and Ali, 2022). We can say that the photooxidation lifetime of methylamine to the sink of CH2NH2 is nearly 13-14 h. Based on current data and previous results, we can say that the $CH_3NH_2 + OH/O_2 \rightarrow CH_2O + HO_2$ via ${}^{\bullet}CH_2NH_2$ leading to the formation of CH₂NH is both kinetically and thermodynamically more favorable than the $CH_3NH_2 + OH/O_2 \rightarrow CH_3NH + HO_2$ via CH_3NH^{\bullet} under tropospheric conditions. This result is also consistent with its isoelectronic analogous (Ali et al., 2019) reaction CH₂OH + OH/O₂ → CH₂O+ HO₂. Generally, the effective rate coefficients for the role of water and ammonia reactions are smaller than those of the free ^oCH₂NH₂ + O₂ reaction system in the temperature range of 200 K-400 K. Therefore, the effect of CH2NH2 + O2 with H2O/ NH₃ is less important for the sink of CH₂NH₂ in a gas-phase atmospheric reaction.

To understand the impact of INT1 in budget calculations, we have calculated the atmospheric lifetime of \sim 3 microseconds of $^{\circ}$ CH₂NH₂ with its reaction O₂, indicating that the formation of

CH₂NH is fast under tropospheric conditions (i.e., at 225 K and an altitude of ~10–11 km) when taking an average concentration of O_2 radicals in the upper troposphere of ~1 × 10^{16} molecule cm⁻³. It is of interest to know whether HO_2 ····CH₂NH₂····H₂O can be produced from the reaction of CH_2NH ····H₂O + O_2 reaction under atmospheric conditions. For this purpose, we calculated the pseudo-first-order rate coefficients of decay of ${}^{\bullet}CH_2NH_2$ ····H₂O+ O_2 at 300 K using concentration $[O_2] = 1 \times 10^{16}$ molecules cm⁻³. The decay rate of INT2h producing CH_2NH_2 +H₂O-HO₂ was found to be 8 × 10^{-3} s⁻¹, which led to the 2 min of the lifetime of HO_2 ····CH₂NH₂····H₂O. Therefore, we can say that under tropospheric conditions, HO_2 ····CH₂NH₂····H₂O can be produced from the reaction of ${}^{\bullet}CH_2NH_2$ + O_2 (+H₂O).

To understand the fate of the CH₃NH[•] radical with its reaction with O2 under the PES (see Supplementary Figure S8), rate coefficients for the addition of O2 to CH3NH are investigated using the RRKM/ME simulation. The O2 radical mildly reacts with CH₃NH with the ~6 kcal/mol below the reactants. Unlike the formation of common aminoperoxy radicals, O2 addition to CH₃NH proceeds with a transition state TS1a with ~5 kcal/mol of barriers, leading to the formation of Int2 and dissected to Int-2 via a five-membered ring transition state with the barrier height of ~13 kcal/mol (with respect to Int-ad) and dissociated with barrierless process to CH2NH + HO2. We predicted the rate coefficients using the direct reaction, i.e., CH₃NH + O₂→TS2a→CH₂NH + HO₂ (Supplementary Figure S8, in blue), CH_3NH^{\bullet} the indirect reaction, i.e., \rightarrow TS1a \rightarrow CH₃NHOO \rightarrow TS2a \rightarrow Int2 \rightarrow CH₂NH HO_2 (Supplementary Figure S8, in black). The calculated rate coefficient for the direct formation of $CH_2NH + HO_2$ is 7×10^{-16} cm³ molecule⁻¹ s⁻¹, which is at least two orders of magnitude smaller than the indirect reaction. Based on our ME calculation, we can say that the formation of CH₂NH does not come from the CH₃NH +O₂ reaction because the reaction is quite slow under tropospheric conditions.

The mechanistic and kinetic analysis suggests the formation of these two *CH2NH2 and CH3NH*, and our overall understanding of atmospheric and interstellar oxygen chemistry remained uncertain. Although many experimental and computational efforts over the past decade on reaction rate coefficients and branching ratios have been made, our knowledge of the chemical pathways theorized for the O2 reaction in two different environments is not clear. Therefore, as suggested in a previous study, at a low temperature, i.e., <100 K, the formation of CH₃NH is dominated over the formation of CH2NH2 under an interstellar cold medium (Gonza'lez et al., 2022). We have performed the RRKM/ME simulation at below 100 K and found that the formation of CH₃NOO is dominated under high-pressure limit conditions, i.e., >10 bar and below 100 K. Under low pressure and low temperature, the reaction goes back to the reactant. The formation of CH3NOO in the ISM medium is not clear, but our analysis suggests that the reaction CH₃NH[•]+O₂ does nothing under ISM conditions. We have also carried out our RRKM/ME simulation under combustion conditions (>1000 K and HPL), and our analysis suggests that the formation of CH₃NOO and CH₂NH is even negligible. It may also be suggested that such a reaction might take place in a more polluted situation.



There has been considerable speculation about what will happen after the formation of CH₂NH (Rissanen et al., 2014; Ali et al., 2018; Ali, 2020). Figure 15 shows the atmospheric degradation reaction of CH₂NH with various possible atmospheric species. The result in the figure shown is based on our previous calculation, except for the reaction of CH₃NH[•] + O₂, which is re-calculated. As shown in Figure 15, we can see that the reaction with water does not lead to the formation of NH₃ and CH₂O as suggested in our previous work under tropospheric conditions (Ali, 2020). We also tried to find out if the backward reaction was favorable; for that, we have set up an ME simulation and predicted that this reaction would do nothing in the troposphere (Ali et al., 2016). We also calculated the rate coefficients for the reaction of CH2NH + OH radicals, and the mechanism has already been discussed in our previous work (Ali and Barker, 2015). The results show the formation of CH₂N°+ and °CHNH as major products and show a similarity between isoelectronic analogous systems, i.e., CH₂O and CH₂CH₂. As suggested in Rissanen et al. (2014) and Ashraful and Silva (2020), water may favor the formation of CH₂N and CHNH; therefore, we have also analyzed the effect of water molecules on the CH₂NH + OH reaction (Ali et al., 2019). We have found that the reaction rate coefficients increase when the concentration of water molecules is not included in the calculation and decrease in the presence of water (Ali et al., 2019). Based on our current and previous findings, we propose that the formation of HCN could be the major product when O₂ radicals react with *CHNH radicals. We also suggest that the gasphase formation of CH_2NH from the $CH_2N^{\bullet} + O_2$ reaction will be even slower, as shown in the previous degradation mechanism (see Figure 14). Again, this is due to less favorable N—O-O bond formation than C-O-O bond formation. We can also suggest that the formation of HCN in its presence may be unimportant under tropospheric conditions. Our calculation suggests that oxidation pathways may also contribute to the HO_x abundance under the tropospheric conditions, as shown in the degradation reaction mechanism (Figure 15 and Figure 14). Such state-of-the-art kinetics provides a clue to the formation of HCN under tropospheric conditions. Our result may be helpful in setting up an experimental analysis for the formation of HCN under tropospheric conditions.

4 Conclusion

In this work, the rate coefficients and branching fraction for $^{\bullet}$ CH₂NH₂ + O₂, $^{\bullet}$ CH₂NH₂ + O₂ (+H₂O), and $^{\bullet}$ CH₂NH₂ + O₂ (+NH₃) for the formation of methanimine (CH₂NH) and HO₂ have been investigated using CCSD(T)//M06-2X/6-311++G (3df, 3pd) coupled with the RRKM/ME simulation. The results show that CH2NH2 + O2 leads to the formation of CH₂NH at temperatures <300 K and goes back to reactants (*CH₂OH + O₂) at high temperatures (>300 K). When the water/ammonia molecule is added to the CH2NH2 + O2 reaction, it favors the formation of CH2NH at a temperature <300 K. The NH₃- and H₂O- assisted rate coefficients are at least 1010-1012 (Gonza'lez et al., 2022) and 106 times, respectively, smaller than those of the free reaction; thus, we can say that the effect of NH₃/H₂O on °CH₂NH₂ + O₂ has less importance in the troposphere. Under tropospheric conditions, the reaction CH₃NH₂ + OH/O₂→ CH₂NH + HO₂ via CH2NH2 leading to form CH2NH + HO2 is both kinetically and thermodynamically more favorable than reaction CH₃NH₂ + OH/O₂ \rightarrow CH₂NH + HO₂ via $^{\bullet}$ CH₃NH. The mechanism indicates that a single NH₃/H₂O molecule has the potential to increase the branching fraction in a gas-phase reaction at a lower temperature <300 K and slower reaction at a higher temperature. Such results are promising, and chemical kinetic data can be beneficial for the future implementation of ammonolysis and hydrolysis of other carbon-centered hydroxyl compounds. In previous studies, researchers stated that the reaction CH2NH may be favorable in water; our study demonstrated that water increases the formation of CH2NH. Our results also indicate the formation of HCN may have come from the reaction going via a carbon-centered radical instead of an N-centered radical. Experimental analysis is required to validate this finding. Such chemical kinetic analysis is interesting; chemical kinetics details can be useful to understand the bigger amine/imine that may lead to the formation of HCN, and N2O may increase the reaction rate.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material; further inquiries can be directed to the corresponding author.

Author contributions

MD and MA completed all the electronic structure calculations, and MA completed the chemical kinetic calculations. MD and MA prepared the draft of the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2023.1243235/full#supplementary-material

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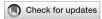
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Experimental study on flow resistance characteristics of the uniform steam injection outflow control device

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Introduction: Multi-point steam injection technology is a new completion method for heavy oil horizontal wells to solve the uneven distribution of the intake profile in the horizontal section. It is equipped with the flow control device to achieve the effect of balanced steam injection.

Methods: The steady-state experiment method was adopted; Considering the variable mass complex flow of the steam—liquid two-phase flow in the downhole flow device, the pressure loss of downhole tools through uniform steam injection with different steam—liquid compositions was tested, the influencing factors of the pressure drop were analyzed, and a more reliable pressure drop calculation method was established.

Results: The overflow pressure drop can be adjusted by changing the aperture, steam dryness, and fluid flow of the downhole outflow control device (OCD).

Discussion: By comparing the experimental and theoretical results, the calculation method of the overflow resistance of single-phase and steam-liquid two-phase fluids in OCD is given, and the error is within the usable range.

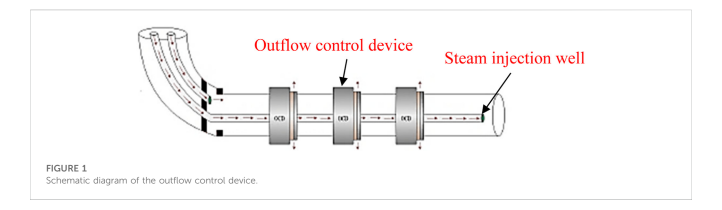
KEYWORDS

outflow control device, two-phase fluids, uniform steam injection, flow resistance characteristics, pressure drop

1 Introduction

In the process of steam injection in long horizontal wells, due to reservoir heterogeneity, heel—toe effect, and natural fractures in the reservoir, the suction profile of steam injection wells is difficult to maintain a balanced development, and steam channeling is easy to occur, and the heel end, high permeability interval, and fractures of wells are prone to premature water/steam. Due to low water/vapor viscosity, once water/gas coning occurs in the well, it will quickly form a channel at the coning point, thereby inhibiting oil production elsewhere (Mozaffari et al., 2013).

In order to reduce this non-uniformity, the flow control device, outflow control device (OCD), can be installed at the completion interval to inhibit the flow through the high-speed interval and generate additional pressure drop so as to increase the flow rate of the wellbore interval with high flow resistance, eliminate the non-uniform flow caused by the heel effect of the horizontal well and the non-uniformity of permeability, and ensure the uniform spread of the suction profile along the horizontal well (Parappilly, and Zhao, 2009; Gu et al., 2014a; Dong et al., 2014b). The combination of a packer and OCD is usually used in production, which can achieve uniform oil drainage in heterogeneous reservoirs and



maintain liquid output balance by limiting different oil production indexes in each section so as to delay bottom water coning, prolong anhydrous or bottom water oil production period, and improve oil and gas well production and recovery efficiency (Sun et al., 2018a; Sun et al., 2019). It is an advanced measure to stabilize oil and water and control water.

At present, a variety of OCDs has been developed in China. According to the mechanism of action, it can be divided into nozzle, spiral channel, and nozzle types (Huang et al., 2018a). The current limiting mechanism of nozzle-type OCD is that the throttle pressure drop is generated by the contraction of the flow channel when the fluid passes through the device (Kumar et al., 2010; Huang et al., 2018b). The spiral-channel-type OCD wraps one or more flow channels around the tubing to generate an additional pressure drop using friction (Rivas and Gates, 2018; Dong et al., 2020a; Dong et al., 2020b). Unlike the instantaneous pressure drop produced by the nozzle, this design produces a segmented pressure drop over a relatively long area, which is more resistant to the erosion and clogging of fluid particles during the drilling fluid cycle (Dong et al., 2020a). The nozzle-type OCD has a limited flow in a long nozzle. Compared with the nozzle-type OCD, owing to its longer spray irrigation, the resistance loss along the path is larger and the local resistance loss is smaller at the same strength (Hasan and Kabir, 1994; Hasan et al., 1998; Sun et al., 2018b). Therefore, compared with the three OCDs, the nozzle-type OCD is more resistant to the erosion and blockage of fluid particles (Hasan and Kabir, 2005; Caetano, 1985). Compared with the spiral-channel-type OCD, the resistance loss along the nozzletype OCD plays a minor role in the pressure drop composition of the nozzle-type OCD, so the nozzle-type OCD is less sensitive to viscosity (Caetano et al., 1992; Gu et al., 2014b; Gu et al., 2015). The installation position of OCD on the horizontal steam injection well is shown in Figure 1.

Aiming at the completion characteristics of horizontal well OCD, this paper established a mathematical model of horizontal well variable mass flow under the condition of OCD installation based on the equivalent diameter model of the target well section, heterogeneous reservoir skin factor model, horizontal wellbore variable mass flow pressure drop model, and OCD pressure drop model by applying the principle of potential superposition. The effects of OCD aperture, steam dryness, and steam injection speed on the pressure drop were compared, and the OCD structure parameters and steam injection parameters were optimized. This

method provides a new optimal control method for horizontal well uniform steam injection (Yin et al., 2019; Liu et al., 2022; Yin and Linga, 2019) and offers technical support for OCD completion optimization design and oil increase and water control in the horizontal well.

2 Experimental scheme

The main research content of this experiment is to measure the pressure drop of the fluid passing through the downhole flow control device and to establish a set of calculation methods of flow control resistance through the study of fluid resistance law so as to determine a reasonable OCD design method to achieve the purpose of uniform steam injection (Least et al., 2013; Yu et al., 2010). In order to study and analyze the flow resistance of different components of fluid that passes through the flow control device and analyze the influence of gas-liquid composition on the pressure drop, the pressure drop of single-phase liquid passing through the control device was first tested, and the pressure drop of fluid passing through the control device was analyzed by changing the flow rate, hole diameter, and fluid flow rate of the control device (Chesney et al., 2015). Then, the influence of the mixture flow rate and mass gas content on the pressure drop was analyzed by changing the mass gas content (Temizel et al., 2019).

2.1 Experimental apparatus and materials

The downhole tool flow resistance test system mainly comprises five parts: the gas supply system, liquid supply system, outflow control device (OCD) simulation system, measurement system (Gai et al., 2010), and data acquisition system (Figure 2). The fluid supply system consists of a liquid pump, inverter, air compressor, and gas pressure-reducing valve (Luo et al., 2015). The liquid pump uses a non-pulse screw pump. Its head measures 180 m, and the displacement range is 0–45 m³/h (Wang et al., 2016); the maximum pressure of the air compressor is 1.8 MPa, the maximum displacement is 5.1 m³/min, and the noise is 56 db; the gas storage tank volume is 1.5 m³, and working pressure is 0.8 MPa; it is made of Q345 material, equipped with a 1 MPa safety valve, and facilitates spray treatment. The gas-reducing valve is of YK43X type, with the maximum pressure of 1.6 MPa and output pressure range of

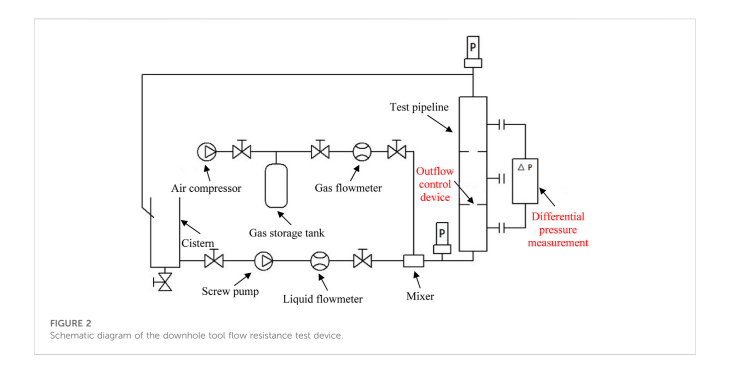




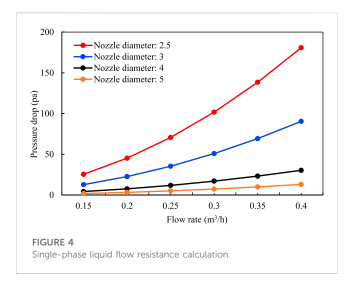
FIGURE 3
Flow control device simulator

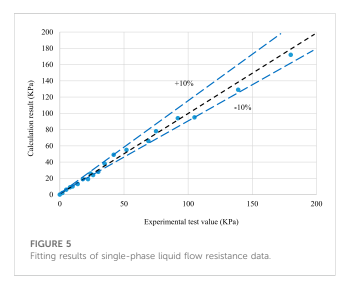
0.1–1 MPa. The accuracy of the pressure transmitter is 0.25% F.s, and the measuring ranges are 0–5 MPa and 0–1.6 MPa, respectively (Prakasa et al., 2019). The accuracy of the temperature sensor is class A, with a measuring range of 0°C–150°C, the flow sensor accuracy of 0.2%–0.5%, liquid flowmeter range of 0–16 m³/h, and gas flowmeter of 0–5.0 m³/min. The sensor can communicate with the computer through the A/D conversion board. The nozzle-type OCD is installed on the test completion string. The data acquisition and processing system comprises digital acquisition card, computer, and software. It carries out real-time data acquisition and post-processing and prepares original data reports, analysis reports, and curves, creates output database files, indicates the real-time display of control elements' working status, and displays and prompts users to each stage of the workflow and pressure upper limit alarm.

2.2 Test procedure

Due to the nozzle-type flow control device used in the FluxRite[™] completion string in the Mackay River oil sand, the fluid pressure drop through the flow control device was studied only under the conditions of single-phase liquid and high gas–liquid ratios (dryness).

First, eight sizes of simulated nozzles were made according to the nozzle sizes of the FluxRite $^{\text{TM}}$ flow control device (2.5 mm, 4.0 mm), and four of them (2.5 mm, 3.0 mm, 4.0 mm, and 5.0 mm) were selected to conduct pressure difference loss experiments on single-phase liquid flow resistance (Figure 3). The test medium was tap water, with reference to the hot water injection flow of horizontal wells in Canada's Mackay River block (0.3 m³/h), and the flow rates were 0.15 m³/h, 0.2 m³/h, 0.25 m³/h, 0.3 m³/h, 0.35 m³/h, and 0.4 m³/h. The pressure drop of the fluid through the simulated



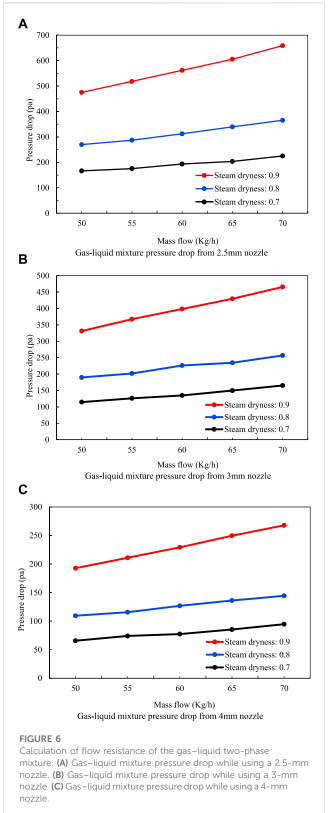


nozzle was recorded by the data acquisition system, and the relationship between the pressure drop and flow rate was analyzed.

Then, three nozzles of sizes 2.5 mm, 3.0 mm, and 4.0 mm were selected to carry out the resistance characteristic experiment of the gas-liquid two-phase flow through simulated nozzles. The gas-liquid mixture consisted of air and water, with reference to the flow rate of wet steam injection in horizontal wells in Canada's Mackay River block (60 kg/h), and its mass flow rate was 50 kg/h, 55 kg/h, 60 kg/h, 65 kg/h, and 70 kg/h. Considering that the steam-assisted gravity drainage (SAGD) steam injection process is a relatively high-dryness flow process, the dryness of the gas-liquid mixture (that is, the mass content of the gas in the mixture) was set to 0.9, 0.8, and 0.7. The pressure difference of the fluid through the simulated nozzle is recorded by the data acquisition system, and the relationship between the pressure drop and the flow rate and the dryness of the mixture is analyzed.

2.3 Experimental procedure

The experimental procedure is as follows:



(1) Screw pump pumps the fluid in the liquid storage tank after suction and pressurization, with a certain flow and pressure through the flow control device.

TABLE 1 Experimental data of single-phase liquid flow resistance (kPa).

Flow rate (m³/h)		0.15	0.2	0.25	0.3	0.35	0.4
Nozzle diameter/mm	2.5	25.42	45.20	70.62	101.70	138.42	180.80
	3	12.72	22.61	35.32	50.87	69.23	90.43
	4	4.26	7.58	11.84	17.05	23.20	30.31
	5	1.83	3.25	5.07	7.30	9.94	12.98

TABLE 2 Pressure difference data of the gas-liquid mixture flow resistance through the 2.5-mm simulated nozzle.

Mass flow (kg/h)		50	55	60	65	70
Dryness	0.9	475.0	518.4	561.8	605.3	658.8
	0.8	269.9	287.1	312.3	339.6	365.4
	0.7	166.4	175.4	193.5	203.5	225.1

- (2) The liquid flow rate is measured by the liquid flowmeter, the size of the flow rate is controlled by the motor frequency converter, and the flow rate of the valve is recorded in real time.
- (3) The pressure drop of the liquid through the flow control device is reflected and recorded in computer software by the differential pressure meter connected at both ends.
- (4) The flow rate and pressure drop are monitored by the computer in real time, such that the pressure drop generated when fluid passes through the control device under a certain flow condition can be obtained.
- (5) In the process of a single-phase liquid experiment, the liquid can be returned to the liquid storage tank through the pipeline for recycling.
- (6) After the completion of the single-phase liquid experiment, start the compressor, adjust the gas-phase flow rate and liquid-phase flow rate, and when the flow is stable, record the pressure drop and dryness of the gas-liquid mixture in the outflow control device.

3 Results and discussion

3.1 Analysis of single-phase liquid flow resistance

The pressure difference data of single-phase liquid flow resistance through the simulated nozzle are shown in Table 1 and the pressure drop curves under different nozzle diameters are shown in Figure 4. In the general literature, the formula for calculating the pressure drop of the fluid through the orifice plate or nozzle is used to obtain the throttle pressure difference when the fluid flows through the control valve:

$$\Delta p = \frac{1}{2} \left(\frac{G_l}{C_d A_p \rho_l} \right)^2 \rho_l,\tag{1}$$

where Δp is the pressure difference before and after the throttle hole of the control valve, that is, the throttle pressure drop in Pa; $A_{\rm p}$

TABLE 3 Pressure difference data of the gas-liquid mixture flow resistance through the 3.0-mm simulated nozzle.

Mass flow (kg/h)		50	55	60	65	70
Dryness	0.9	331.2	367.3	398.4	429.5	465.6
	0.8	189.7	201.8	226.2	234.5	256.7
	0.7	114.6	126.1	134.9	149.7	165.2

TABLE 4 Pressure difference data of the gas-liquid mixture flow resistance through the 4.0-mm simulated nozzle.

Mass flow (kg/h)		50	55	60	65	70
Dryness	0.9	192.8	211.0	229.3	249.6	267.9
	0.8	109.5	115.7	126.9	136.2	144.4
	0.7	65.7	74.1	77.4	85.4	94.7

is the control valve orifice area in m^2 ; and C_d is the flow coefficient without dimensionality. G_I is the steam mass flow rate in kg/s, and ρ_I is the fluid density in kg/m³. The value of C_d in common flow control valves ranges from 0.66 to 1.

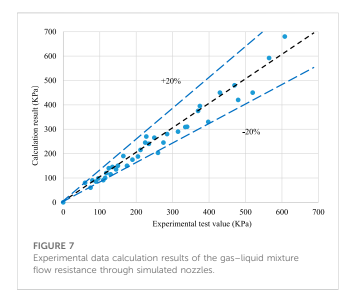
According to the method of drawing the nozzle loss curve, the flow coefficient $C_{\rm d}$ of the fluid passing through the nozzle should not be a constant but a function of the hole diameter. According to the data of the pressure drop, flow rate, and hole diameter, the corresponding relationship between the flow coefficient $C_{\rm d}$ and hole diameter under different flow rates was obtained, and data fitting was carried out to obtain the fitting relationship between $C_{\rm d}$ and hole diameter:

$$C_d = 1.3052d^{-0.11} = \frac{1.3052}{d^{0.11}}.$$
 (2)

The calculated results of the fitting relationship were compared with the experimental results (Figure 5). The comparison results show that the flow coefficient obtained by fitting can calculate the experimental data more accurately.

3.2 Flow resistance analysis of the gas—liquid two-phase fluid

The Pressure difference data of the gas-liquid mixture flow resistance through the simulated nozzle are shown in Tables 2-4.



The calculation method of the pressure drop of gas-liquid mixture flow through the orifice plate proposed by Chisholm was used for data processing. Through the two-phase flow momentum equation analysis, the pressure drop of the fluid flowing through the orifice plate can be calculated as follows:

$$\Delta p_{\mathrm{TP}}/\Delta p_{\mathrm{LO}} = 1 + \frac{C}{X} + \frac{1}{X^2}, \tag{3}$$

where X^2 is equal to $\Delta p_{\mathrm{LO}}/\Delta p_{\mathrm{GO}}$; Δp_{TP} is the pressure drop of the gas-liquid two-phase mixture through the orifice plate in Pa; Δp_{LO} and Δp_{GO} are the pressure drop when the liquid and gas phases in the mixture, respectively, pass through the orifice plate alone in Pa; and C is a coefficient related to the pressure and slip ratio.

The following formula was used to calculate X^2 :

$$X^2 = \frac{(1-x)^2}{x^2} \frac{\rho_G}{\rho_L}.$$
 (4)

In 1997, Chisholm suggested the following method for calculating the *C* values:

When X < 1, then

$$C = \left(\frac{\rho_{\rm L}}{\rho_{\rm C}}\right)^{1/4} + \left(\frac{\rho_{\rm G}}{\rho_{\rm L}}\right)^{1/4}.\tag{5}$$

When X > 1, then

$$C = \left(\frac{\rho_{\rm m}}{\rho_{\rm G}}\right)^{1/2} + \left(\frac{\rho_{\rm G}}{\rho_{\rm m}}\right)^{1/2}.\tag{6}$$

The calculation method proposed by Chisholm was used to calculate the pressure drop data of the gas-liquid mixture flow through the simulated nozzle. The pressure drop changes under the aperture of 2.5 mm, 3 mm, and 4 mm were obtained, as shown in the Figure 6, and the pressure drop decreased significantly when the nozzle size was increased. The greater the steam dryness, the greater the pressure drop and the better the effect of overflow resistance. The flow rate has little influence on the flow resistance effect, and the flow rate is positively correlated with the pressure drop.

The calculated results were compared with the experimental results (Figure 7). The comparison results show that the calculated values using Chisholm's recommended method were consistent with the experimental results of the gas-liquid mixture, and the calculation error is less than $\pm 20\%$.

3.3 Flow resistance analysis of the gas—liquid two-phase fluid with a large aperture

Since multi-stage OCD completion is usually used in the production process, that is, multiple OCDs are connected in series for steam injection at the same time, this paper designs a variety of large-aperture OCDs to meet production needs and installation on demand. Using the steam dryness of 0.9, steam injection pressure of 2 MPa, large aperture sizes of 20 mm, 50 mm, 80 mm, 110 mm, and 140 mm, the pressure drop and temperature change of steam through a valve under different apertures are calculated; the pressure drop calculation method of the gas-liquid two-phase flow is referred to formulas (3)–(6), and the heat calculation method is as follows:

The temperature and pressure of saturated steam have a coupling relationship, which is given as follows:

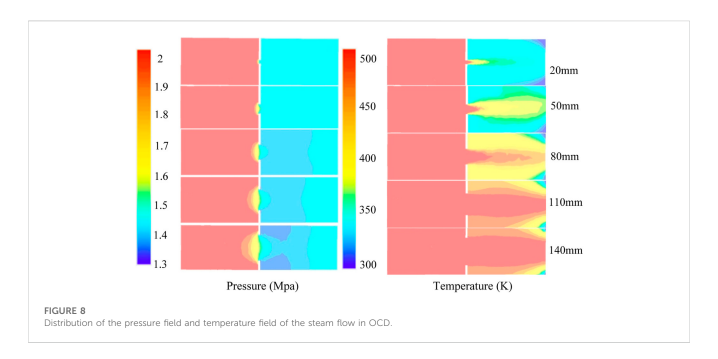
$$T_s = 210.2376 p_s^{0.21} - 30, (7)$$

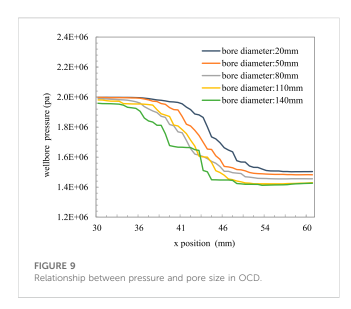
where T_s is the temperature of steam in °C and p_s is the pressure of steam in MPa.

The heat of the injected wet steam will reach the formation along the radial path through the inner tube of the insulated tubing, the insulation layer of the insulated tubing, the outer tube of the insulated tubing, and the annular space between the tubing and casing and the casing and cement ring. In this process, radial heat loss will be generated, which can be bounded by the outer edge of the cement ring. The former is the steady heat transfer, and the latter is the unsteady heat transfer. Convection heat transfer occurs when the wet steam flows through the inner wall of the insulated tubing, heat transfer occurs when it flows from the inner wall of the insulated tubing to the outer wall of the insulated tubing, convection heat transfer and heat radiation occur when it flows in the annular space between the tubing and casing, and heat transfer occurs when it flows in the annular space between the casing and cement ring; hence, the thermal resistance formula is expressed as follows:

$$R = \frac{1}{2\pi h_{1}r_{1}} + \frac{1}{2\pi \lambda_{tub}} \ln\left(\frac{r_{2}}{r_{1}}\right) + \frac{1}{2\pi \lambda_{ins}} \ln\left(\frac{r_{3}}{r_{2}}\right) + \frac{1}{2\pi \lambda_{tub}} \ln\left(\frac{r_{4}}{r_{3}}\right) + \frac{1}{2\pi (h_{c} + h_{r})r_{4}} + \frac{1}{2\pi \lambda_{ras}} \ln\left(\frac{r_{co}}{r_{ci}}\right) + \frac{1}{2\pi \lambda_{rem}} \ln\left(\frac{r_{h}}{r_{co}}\right),$$
(8)

where h_1 is the convective heat transfer coefficient of wet steam in W/(m²·°C) and r1 is the inner tube radius of the insulated tubing in m. λ_{tub} is the thermal conductivity of the inner tube of the heat-insulated tubing in W/(m·°C); r_2 is the inner and outer radius of the insulated tubing in m; λ_{ins} is the thermal conductivity of the insulation layer of the insulated tubing in W/(m·°C); r_3 is the outer tube radius of the insulated tubing in m; r_4 is the outer radius of the insulated tubing in m; h_c is the natural convection heat transfer coefficient in the annular space between the tubing and





casing in W/(m²·°C); h_r is the radiant heat transfer coefficient in the annular space between the tubing and casing in W/(m²·°C); $\lambda_{\rm cas}$ is the thermal conductivity of the casing in W/(m·°C); $r_{\rm co}$ is the outer radius of the casing in m; $r_{\rm ci}$ is the tube radius in m; $\lambda_{\rm cem}$ is the thermal conductivity of the cement ring in W/(m·°C); and $r_{\rm h}$ is the outer radius of the cement ring in m.

The section of steam flowing through the nozzle is suddenly reduced, and the formula for calculating overflow resistance is given as follows:

$$F = \frac{P_1 - P_2}{\gamma} + \frac{\alpha_1 V_1^2}{2g} - \frac{\alpha_2 V_2^2}{2g} - F_{1-2}, \tag{9}$$

where F is the overflow resistance; F_{1-2} is the resistance loss along the path; α_1 and α_2 are the kinetic energy correction coefficients; γ is the fluid weight; and V_1 and V_2 are the average flow rates at the interface.

It can be observed from Figure 8 and Figure 9 that the pressure drop of the gas-liquid two-phase fluid varies greatly under different apertures. Because the mechanism of the local pressure drop of nozzle-type OCD depends on the minimum flow area, with the increase in the aperture, the pressure drop becomes smaller, the temperature range decreases, the heat transfer speed increases, and the heat loss decreases. When the pore diameter is greater than 110 mm, the effect of the steam pressure drop through the pore is not obvious, and the effect of OCD current limiting is worse.

4 Conclusion

- (1) A simulation experimental device was established to carry out the flow resistance experiment of the single-phase liquid and gas-liquid mixture through the nozzle/orifice plate flow control device simulation system.
- (2) Data analysis shows that the experimental data of the single-phase liquid flow can reliably be predicted using the formula of the liquid pressure drop through the orifice plate and the flow coefficient obtained by fitting the experimental data, and the prediction accuracy is less than ±10%; the calculation error of the experimental data of the gas-liquid mixture can be less than ±20% by using the method proposed by Chisholm.
- (3) The factors affecting the flow resistance effect of the gas-liquid two-phase flow were compared and analyzed, and the overflow pressure drop effect under different pore sizes, steam dryness, and steam flow rates was simulated. It was concluded that the pore size had a significant effect on the pressure drop effect, and the pressure drop amplitude became smaller with the increase in the pore size. In order to achieve an obvious throttling effect, the maximum pore size should not exceed 80 mm; the higher the steam dryness, the more obvious the pressure drop effect is. The flow rate has little influence on the overflow resistance, and the

overflow pressure difference becomes larger with the increase in the flow rate.

Data availability statement

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

Author contributions

QD: conceptualization, data curation, formal analysis, investigation, methodology, project administration, resources, software, supervision, validation, visualization, writing-original draft, and writing-review and editing. ML: methodology, resources, and writing-review and editing. CL: data curation, supervision, and writing-review and editing.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Fractionation of *Xanthium* strumarium L. foliage phenolics, in-vitro antioxidant activities, and in-vivo anti-diabetic potential

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Introduction: The present research aimed to fractionate *Xanthium strumarium* L. (XSL) foliage phenolics into a set of solvents and evaluate their antioxidant potential and *in-vivo* anti-diabetic activity against Alloxan monohydrate-induced diabetic mice.

Methodology: For this purpose, XSL foliage was fractionated into petroleum ether, ethyl acetate, ethanol, and water *via* orbital type shaking and tested for the presence of phenolics, and their antioxidant and antidiabetic potential.

Results and discussion: The results revealed that the ethyl acetate fraction of XSL foliage contained the highest amount of total phenolics 95.25 mg GAE/g of extract, followed by ethanol (65.14 mg GAE/g), petroleum ether (25.12 mg GAE/g), water (12.20 mg GAE/g), and XSL powder (69.13 mg GAE/g). At the end of treatment time (day 18 of oral administration of 400 mg/kg body weight of mice), the ethyl acetate fraction significantly ($p \le 0.05$) lowered blood glucose level (353 \pm 10.6 to 220 \pm 25.5 mg/dL) which might due to the elevated level of phenolic compounds in this fraction.

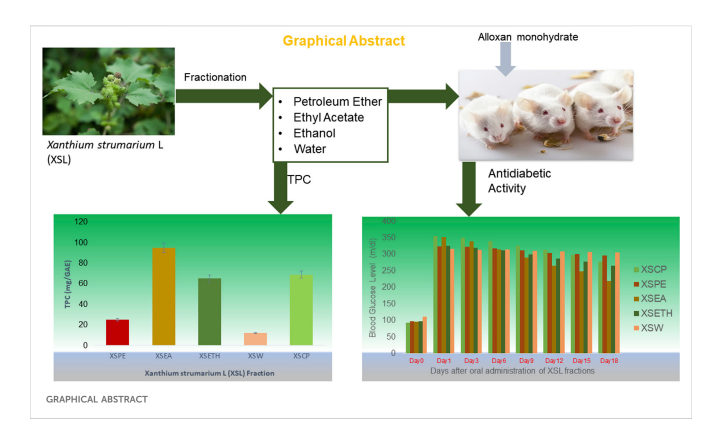
Conclusion: Overall, it can be speculated that ethyl acetate and ethanol may work efficiently for the enrichment of XSL phenolic without compromising their antidiabetic potential.

KEYWORDS

Xanthium strumarium, L. Foliage, fractionations, phenolics, antioxidants, antidiabetics

1 Introduction

Despite the prevalence and widespread use of anti-diabetic medications like metformin, sulfonylureas, thiazolidinediones, and insulin, diabetes ranks as a major global health concern. International Diabetic Federation estimates indicate that the prevalence of diabetes has increased to 463 million in 2019 and figures may rise to 578 (10.2%) and 700 million (10.9%) by 2030 and 2045, respectively (Akhtar et al., 2022). Additionally, it has been predicted that by 2030, diabetes will account for 3.3% of all fatalities worldwide, making it the seventh greatest cause of death (Kokil et al., 2015). For centuries herbs have been used to meet healthcare needs through medical procedures and traditional remedies. Traditional medicine employs at least 1200 different plant species for their potential biological activities and almost half of these have been studied for their phytochemistry, and antioxidant activities (Krupa et al., 2019; Mawoza et al., 2019).



Xanthium strumarium Linn (hereafter XSL) belongs to the Asteraceae family and is commonly identified as cocklebur, sheep bur, hedgehog bur weed, ditch bur, sea burdock, clot bur, and button bur by different civilizations in the world (Ghahari et al., 2017). The plant has habitats worldwide, but temperate zones are the areas of high prevalence, especially India, Russia, Iran, Australia, North Korea, Japan, Pakistan, America, South Africa, China, Eastern Asia, and some parts of South Asia. It frequently grows along roadsides, and on plains, hills, and mountains. The flowering period starts during July/ August whereas fruits are usually ripe in September/October (Fan et al., 2019). The fruit as a whole clings to animal fur for its dispersal and is considered poisonous, however, it possesses cytotoxic, sedative diuretic, antitussive antimalarial, antispasmodic, anti-rheumatic antibacterial, and antifungal properties (Clayton et al., 2020). Few researchers have evaluated the antidiabetic potential of X. strumarium L. (XSL), but the majority of these studies utilized aqueous or alcoholic extracts of XSL foliage (Ahmad et al., 2016). An acute review of previously published research regarding phenolics reveals that the solubility of these compounds varies widely and similar can be speculated about the stability and biological functions of these bioactives. Keeping in view these facts, we have planned to fractionate XSL foliage phenolics into a range of solvents including petroleum ether, ethyl acetate, ethanol, and water and each fraction of XSL foliage was subsequently assessed for its total phenolic content (TPC), radical scavenging capacity, in-vitro antioxidant activity, and antidiabetic potential (in-vivo).

2 Materials and methods

The experimental work regarding the fractionation of XSL foliage was done in the Biochemistry laboratory of Government

College University Lahore. The *in-vivo* trials associated with the present research were carried out at the animal house in the Lahore College for Women University's Zoology Department. Formal approval was obtained from the institutional ethical committee (No. GCU-IIB-2596) dated 31 October 2022. Animals (Swiss albino mice) were supplied by F. Z traders, Lahore, Pakistan, and at the time of the experiment the average age and weight ranged from 4 to 8 weeks and 22–24 g, respectively. The chemicals and solvents used during the current study were of analytical grade and acquired from the companies Sigma Aldrich Chemical Co. (United States) and Merck (Germany).

2.1 Sample collection

The XSL foliage was collected during the month of June 2022 from a remote location in Pakistan's District Narowal and was identified as rough cocklebur (voucher no. LCW -1016) by Dr. Shubnam Shaheen Associate Professor, Department of Botany, Lahore College for Women University Lahore.

2.2 Preparation of XSL fractions

The XSL foliage was air dried at 40°C after being cleaned with distilled water to remove dirt and contaminants. The dried XSL foliage was powdered prior to the fractionation with particle mesh sizes ranging from 24 to 60 sorted by electromagnetic vibrator for 15 min, packed in an opaque plastic bag, and stored. Petroleum ether, ethyl acetate, ethanol, and water were

sequentially used as solvents during the fractionation process. A carefully-weighed 25 g of dried XSL foliage powder was placed in a 500 mL Erlenmeyer flask and shaken with 250 mL of petroleum ether for 12 h in an orbital shaker (Gallenkamp, UK) at 250 rpm under ambient conditions 25°C-30°C. The resultant aliquot was filtered using Whatman filter paper 1, and residues were mixed with the second extraction solvent, i.e., ethyl acetate, and shaken under the above-mentioned conditions. The residues of step 2 and step 3 were subsequently subjected to ethanol- and water-based fractionation following the above-mentioned procedure. All four fractions were separately dried using a rotating evaporator (SB-651; EYELA, Tokyo, Japan) operated at 35°C and reduced pressure, weighed to calculate the percent yield of each fraction (g/100 g of XSL powder), and stored at -10°C until the required dosages were made. Finally, the amount of extractable bioactives (total extract) was calculated by adding the percent yield of each fraction.

2.3 Total phenolic content

The TPC in every fraction of XSL foliage powder was measured using Folin-Ciocalteu reagent as documented by (Elagdi et al., 2023). Gallic acid was processed as a positive control to express the results as milligrams of gallic acid equivalent (mg GAE)/g) of extract. The five different concentrations of the gallic acid standard (50, 40, 30, 20, and 10 mg/mL) and 1.0 mg/mL of each XSL foliage powder fraction were introduced into the test tube containing 2.5 mL of 10% (v/v). To the Folin-Ciocalteu reagent was added 2 mL of 7.5% (w/v) $\rm Na_2CO_{3}$, and the mixture was allowed to stand in the dark at room temperature for half an hour. The absorbance was measured at 760 nm using a Shimadzu 160-UV spectrophotometer. For each extract, all the calculations were done in triplicates.

2.4 DPPH assay

The radical scavenging potential of each XSL foliage fraction was estimated by using the method described by Villano et al. with minor changes (Villaño et al., 2007). In this assay, 0.1 mM solution of 1,1diphenyl-2-picrylhydrazyl and 1.0 mg/mL of each XSL extract was prepared in HPLC grade methanol. The equal volumes 2.5 mL of DPPH solution and extract were incubated at room temperature for half an hour under dark, and at 517 nm absorbance was measured and compared with ascorbic acid acting as a standard. The free radical scavenging potential of the XSL fraction was expressed as %inhibition calculated by the following formula where As and Ac stand for the absorbance of the sample and control respectively. The IC₅₀ (Conc. providing 50% inhibition) value is used to express the DPPH assay results. IC₅₀ value can be measured by plotting a graph between the scavenging effect and the corresponding extract concentration. AAI (Antioxidant activity index) is used to measure the antioxidant potential and can be calculated by dividing the final DPPH concentration (µg/ mL) by IC₅₀ (μg/mL) (Scherer and Godoy, 2014).

 $%Inhibition = [(Ac-As)/Ac] \times 100$

2.5 FRAP assay

The ferric-reducing ability of plasma (FRAP) comprising XSL foliage fractions was determined by using the strain and Benzie method with minor changes (Benzie and Strain, 1996). In this method, 300 μL distilled water was mixed in 100 μL of plant extract followed by the addition of 3 mL of frap reagent. The frap reagent was prepared by mixing 25 mL acetate buffer (0.3 M) at PH = 3.6 and 2.5 mL (10 mM) of TPTZ (2, 4, 6- tripyridyl-s-triazine) in hydrochloric acid (40 mM) and 2.5 mL of FeCl₃ (20 mM). The mixture was incubated under ambient conditions (30°C) and absorbance was measured at 593 nm by a Shimadzu 160-UV spectrophotometer. Frap assay evaluates the antioxidant activity of plant extract by reducing metal ions using electron donation. Ascorbic acid was used as the standard and the FRAP value was measured by the following equation, Where $A_{\rm O}$ and $A_{\rm S}$ are the absorbance of the standard and sample respectively.

Frap value $((\mu mascorbate/g) = As-Ao/Ao$

2.6 Induction of diabetes in mice

For the animal study, 45 male albino mice were selected and were kept for 1 week of acclimatization. With a maximum of six animals per cage, the animals were housed in sterile polypropylene cages and kept at room temperature. The bedding material was sterile rice husk. These animals were housed in a controlled setting with respect to temperature (25°C), humidity (45%–57% range), and light period (12: 12-h darklight cycle). Alloxan monohydrate was intraperitoneally injected at a dosage of 120 mg/kg to cause hyperglycemia in male albino mice that were more than 8 weeks old and weighed 22–24 g. The blood glucose levels were assessed by tail clip sampling 48 h following alloxan administration. When the blood sugar level exceeded 185 mg/dL, mice were considered diabetic (Nagappa et al., 2003). The animals were divided into seven groups each having six mice and the body weight and blood glucose (using glucometer) level of control and treated groups were monitored on days 1, 3, 6, 9, 12, 15, and 18 of treatment.

C: Mice who had not been given any drug or alloxan monohydrate (Normal control).

DC: Mice were given alloxan monohydrate (Diabetic control). XSCP: Diabetic mice were administered with crude powder of XSL.

XSPE: Diabetic mice were administered a petroleum ether fraction of XSL.

XSEA: Diabetic mice were administered with ethyl acetate fraction of XSL.

XSETH: Diabetic mice were administered an ethanolic fraction of XSL.

XSW: Diabetic mice were administered with an aqueous fraction of XSI.

2.7 Acute toxicity studies

The process was completed in accordance with OECD (Organization for Economic Co-operation and Development)

TABLE 1 The XSL foliage fraction yield in various solvents and their Total Phenolic Content (TPC).

Fractions	% yield (w/w)	TPC (mg GAE/g)
XSPE ¹	0.033 ± 0.01 ^a	25.12 ± 7.06^{ab}
XSEA ²	0.095 ± 0.02^{a}	95.25 ± 7.41°
XSETH ³	0.301 ± 0.07 ^b	65.14 ± 7.06 ^b
XSW ⁴	0.402 ± 0.28 ^{bc}	12.2 ± 5.80°
Total	$0.828 \pm 0.38^{\rm d}$	197.71 ± 27.33 ^d
Crude powder (XSCP ⁵)		69.13 ± 6.01^{bc}

The values are mean \pm SD, of triplicate experiments, the different superscript letters over each range denote the significantly different ($p \le 0.05$) values.

guidelines to check the toxicity (Ahmad et al., 2016). To investigate the acute toxicity, different doses of XSL foliage fractions were administered to male Swiss albino mice. The control group received the vehicle (normal saline only), whereas the treatment groups orally received plant fractions in dosages of 2 and 5 g/kg. The adverse effects were thoroughly monitored for 2 days after the dose was introduced. The body weight of the mice before and after administration, any sign of toxicity such as changes in the fur, skin, and eyes, as well as changes in the respiratory, circulatory, and central nervous systems, behavior patterns, signs of tumors, salivation, diarrhea, sleep, and coma were also noted.

2.8 Blood collection and determination of blood glucose

To determine the blood glucose level blood samples were collected from fasted mice on days 0, 1, 3, 6, 9, 12, 15, and 18 of the treatment. Blood was collected¹ from the mouse tails by snipping with a sharp razor, and glucose level was measured by glucometer (Accu-Chek Active, Roche Limited, Pakistan)² as described by Arya et al. (2012).

2.9 Statistical analysis

All the parameters including extract yield, TPC, and DPPH radical scavenging activity were measured in triplicate and the estimates were compared³ for significant difference $p \le 0.05$ by using Duncan's multiple range⁴ test (MRT) (Tallarida and Murray, 1987).

3 Results and discussion

Table 1 shows the percentage yield⁵ of fractions of XSL foliage depending on the solvent polarity that significantly affected the fraction yield in the order of petroleum ether < ethyl acetate < ethanol < water. Fraction yield depends upon the solvent polarity

and the difference in fraction yield in different solvents was due to the solubility of phytochemicals in the respective solvent (Dirar et al., 2019). TPC of different fractions of XSL in ethyl acetate was 95.25 ± 7.41 mg GAE/g, followed by ethanol 65.14 ± 7.06 mg GAE/g, petroleum ether 25.12 \pm 7.06 mg GAE/g, and water 12.2 \pm 5.80 mg GAE/g. The TPC in crude powder of XSL foliage was found to be 69.13 ± 6.01 mg GAE/g. In the previous studies TPC value of ethyl acetate and aqueous leave extract of XSL extracted by maceration and hot extraction method was 59.98 and 35.92 mg of GAE/g DW, respectively (Pillai and Thebe, 2023). The TPC value of ethanolic extract prepared from the aerial parts of XSL using the cold percolation method was determined 84.86 ± 5.13 mg of GAE/g DW (Ly et al., 2021). Similarly, aqueous and ethyl acetate fractions were obtained from 80% methanolic extract of XSL dried leaves using the cold percolation method, and their TPC values were determined as 75.24 \pm 13.31 and 166.26 \pm 27.98 mg of GAE/g DW (Guemmaz et al., 2018; Pillai and Thebe, 2023). In other studies ethyl acetate and 80% ethanol extract were extracted from XSL dried leaves by dynamic maceration, static maceration, and soxhlet method, and their TPC values were determined as 70.07 ± 1.6 , 64.51 ± 1.0 , 69.38 ± 1.3 mg GAE/g DW (Ethanol), 27.19 ± 1.0 , 21.98 ± 3.6 and 23.19 ± 0.3 mg GAE/g DW (Ethyl acetate) respectively (Scherer and Godoy, 2014). No literature reported on petroleum ether fraction. The discrepancy in TPC value is due to factors like sample variety, sample quantity, type of extraction techniques, seasonal variation, extracted bioactive components, and geographic location.

3.1 Antioxidant activity

The DPPH assay evaluates the antioxidant's scavenging capacity towards the DPPH radical. DPPH is a stable free radical having a blue color in alcoholic solution, while in the presence of antioxidants its reduction changes color to yellow. DPPH assay depends upon the antioxidant's potential to donate hydrogen or electrons and is spectrophotometrically analyzed. Absorbance is inversely proportional to the antioxidant potential which is directly proportional to % inhibition. Table 2 shows the antioxidant activity of petroleum ether (XSPE), ethyl acetate (XSEA), ethanol (XSETH), and water (XSW) fractions of XSL. The IC₅₀ value decreased in the order of 101.20 > 88.02 > 54.60 > 46.11 and the antioxidant index value (AAI) increased in the order of 1.87 > 0.90 > 0.28 > 0.05 respectively. In the present study the ethyl acetate fraction has the lowest IC₅₀ value followed by the

¹ Petroleum ether fraction of X. strumarium L. Foliage.

² Ethyl acetate fraction of X. strumarium L. Foliage.

³ Ethanol fraction of X. strumarium L. Foliage.

⁴ Water fraction of *X. strumarium* L. Foliage.

⁵ Crude powder of X. strumarium L. Foliage.

TABLE 2 DPPH Radical scavenging activity of different fractions of XSL foliage powder.

Fractions	IC ₅₀ μg/mL	AAI
XSPE	88.02°	0.28 ^b
XSEA	46.11 ^a	1.87 ^d
XSETH	54.60 ^b	0.90°
XSW	101.20 ^d	0.05 ^a

The values are mean \pm SD, of triplicate experiments, the different superscript letters over each range denote the significantly different ($p \le 0.05$) values.

TABLE 3 FRAP Assay of different fractions of XSL foliage powder.

Sample	Absorbance	Frap value (µm ascorbate/g)
XSPE	0.75	0.194 ± 0.04
XSEA	1.19	0.238 ± 0.06
XSETH	0.98	0.212 ± 0.07
XSW	0.59	0.179 ± 0.01

ethanol fraction, while in the previous literature the IC₅₀ value of different solvent extracts of Xanthium strumarium aerial parts has been determined. IC50 values of aqueous and ethyl acetate extracts (obtained by maceration and hot percolation method) have been determined at 2465.21 and 1856.02 µg/mL, respectively (Kim et al., 2005; Guemmaz et al., 2018; Pillai and Thebe, 2023). Similarly Aqueous and ethyl acetate fractions of 80% methanol extract of dried XSL leaves obtained by cold percolation method have demonstrated IC₅₀ values 46.00 \pm 0.0006, 17.00 \pm 0.0004 μ g/mL respectively (Pillai and Thebe, 2023). In another study, 80% ethanol and ethyl acetate extract prepared from XSL dried leaves by dynamic maceration, static maceration, and soxhlet method and their IC₅₀ values were determined as 53.01 \pm 1.20, 47.83 ± 1.40 , $53.34 \pm 1.52 \,\mu\text{g/mL}$ (Ethanol), 369.83 ± 13.58 , 346.35 ± 16.50 , and $423.97 \pm 22.27 \,\mu\text{g/mL}$ (Ethyl acetate) in the DPPH assay (Scherer and Godoy, 2014). There was no literature reported on petroleum ether fraction. The variation in the IC₅₀ in the present study and previous reports might be due to the abovementioned factors as discussed before.

FRAP assay evaluates the antioxidant potential of antioxidants in terms of the reduction of ferric to ferrous ions. Ascorbic acid is used as standard and the values of reducing potential of antioxidants are compared to it. With the increase of antioxidant concentration, absorbance increases, and the resulting reducing power also increases. The frap value will be greater the higher the antioxidant potential. Table 3 depicts the Frap values and absorbance of different fractions of XSL foliage. Ethyl acetate has greater absorbance with FRAP value of 0.238 \pm 0.06 (μ m ascorbate/g) followed by ethanol (0.212 \pm 0.07), petroleum ether (0.194 \pm 0.04) and water (0.179 \pm 0.01) μm ascorbate/g. In the literature cited the ferric reducing capacity of ethyl acetate and water extracts of XSL have been determined to be 0.996 \pm 0.101 and 0.412 \pm 0.009 μm ascorbate/g at a concentration of 100 µg/mL (Kim et al., 2005; Pillai and Thebe, 2023). While the FRAP value in this research work was measured at 30 µg/mL. The FRAP values of ethanol and petroleum ether fractions were not reported in the literature.

3.2 Body weight

Table 4 elaborates on the difference in body weight between the experimental and control group animals after treatment with different fractions of XSL foliage. The mice administered alloxan monohydrate (120 mg/kg) lost body weight from 1% to 3% which was averted by the treatment of different fractions and powdered samples of the selected plant. The control group animals (C) which use only the vehicle shows an increase in body weight from 21.5 \pm 2.12 to 24.75 \pm 1.77 (14%) while diabetic control (DC) loss in body weight from 23.50 \pm 0.71 to 21.05 \pm 0.78 (10%) on day 18 days.

The animals in the XSCP sample lost body weight by 23 ± 2.83 to 22.40 ± 2.69 (3%) on the first day and after receiving with powdered sample increased by 24 ± 2.82 (7%) on the last day of the trial. The XSPE and XSEA animals reduced their body weight from day $1 \ 23.50 \pm 0.71$ to 23.15 ± 0.50 (1%) and 21.50 ± 0.71 to 21.15 ± 0.64 (2%) respectively and after being treated with petroleum ether and ethyl acetate fractions, the weight of XSPE and XSEA animals increased up to 23 ± 0.57 , 24 ± 0.71 (1% and 13%) of their initial body weight. The animals treated with the ethanol and water fractions showed body weight reduction of 23.5 ± 0.71 to 23.05 ± 0.64 (2%), 22.75 ± 1.06 to 22.55 ± 1.06 (1%), and after treatment increased up to 24.15 ± 0.64 , 22.70 ± 0.71 (5% and 1%) respectively. The results shown are based on the mean of six mice and the percentage change indicates the change from day 1 to the end of the study (day 18).

3.3 Antidiabetic activity

The blood glucose level of diabetic mice (CD) was significantly $(p \le 0.05)$ higher as compared to normal mice (C) and the effect of powdered sample and different fractions of XSL on diabetic mice was studied. Data assembled in Table 5 shows the blood glucose levels of the control and treatment groups on days 1, 3, 6, 9, 12, 15, and 18 of treatment. The blood glucose level of C (control group), who simply used a vehicle, did not vary significantly ($p \le 0.05$) during the treatment time (as shown by subscript and superscript letters). However, the blood glucose level of diabetic mice (DC) was significantly ($p \le 0.05$) higher as compared to the control group. Fasting mean blood glucose level of diabetic control DC and XSCP on day one after being diabetic was 345 ± 7.07 mg/dL, and 355 ± 7.07 mg/dL, respectively. On day 18 the blood glucose level of DC (diabetic control) increased up to $364 \pm 2.12 \text{ mg/dL}$, while oral administration of powdered sample to XSCP significantly ($p \le 0.05$) decreased the blood glucose level to 278 ± 10.6 mg/dL (22%). On day one, blood glucose levels of diabetic-induced mice of XSPE, XSEA,

TABLE 4 Effect of different fractions of XSL on body weight in alloxan-induced diabetic mice.

Groups	Doses (mg/kg)		Days after treatment						
		0		3	6	9	12	15	18
С		21.50 ± 2.12	21.65 ± 2.05	21.90 ± 1.98	22.20 ± 1.84	22.55 ± 1.91	22.90 ± 1.84	23.75 ± 1.77	24.75 ± 1.77
DC		23.50 ± 0.71	23.30 ± 0.57	23.25 ± 0.78	22.80 ± 0.85	22.45 ± 0.78	22.00 ± 0.71	21.45 ± 0.78	21.05 ± 0.78
XSCP	400	23.00 ± 2.83	22.40 ± 2.69	22.30 ± 2.69	22.75 ± 2.76	23.05 ± 2.62	23.25 ± 2.62	23.60 ± 2.69	24.00 ± 2.82
XSPE	400	23.50 ± 0.71	23.15 ± 0.50	23.00 ± 0.57	22.50 ± 0.71	22.65 ± 0.64	22.80 ± 0.57	22.90 ± 0.57	23.00 ± 0.57
XSEA	400	21.50 ± 0.71	21.15 ± 0.64	21.60 ± 0.71	21.95 ± 0.64	22.50 ± 0.71	22.85 ± 0.64	23.30 ± 0.71	24.00 ± 0.71
XSETH	400	23.50 ± 0.71	23.05 ± 0.64	22.95 ± 0.64	23.15 ± 0.78	23.55 ± 0.78	23.65 ± 0.78	23.75 ± 0.78	24.15 ± 0.64
XSW	400	22.75 ± 1.06	22.55 ± 1.06	22.30 ± 0.99	22.3 ± 0.71	22.40 ± 0.71	22.45 ± 0.78	22.50 ± 0.71	22.70 ± 0.71

The values are mean \pm SD, of triplicate experiments.

TABLE 5 Blood Glucose Level of alloxan-induced diabetic mice at different intervals after XSL fractions administration.

Groups	Dose (mg/kg)		Days after treatment						
		0		3	6	9	12	15	18
С		103 ± 9.90° a	95.5 ± 0.71° _a	97 ± 4.24° _a	91 ± 1.41° _a	94 ± 2.83° _a	96.5 ± 2.12 ^a _a	93 ± 4.24° _a	97 ± 1.41° _a
CD		105 ± 7.07° a	345 ± 7.07 ^d _d	348 ± 5.66 ^d _d	350 ± 5.66 ^d _d	355 ± 7.07 ^d _d	358 ± 5.66 ^d _d	363 ± 3.54 ^d _d	364 ± 2.12 ^d _d
XSCP	400	93.5 ± 0.71° _a	355 ± 7.07 ^d _d	350 ± 7.07 ^d _d	339 ± 8.49 ° _c	327 ± 9.20 ° _c	313 ± 11.3° _c	299 ± 4.95 ^b _b	278 ± 10.6 b _b
XSPE	400	97.5 ± 3.54° _a	325 ± 21.2 ^d _c	323 ± 21.9 ^{cd} _b	318 ± 21.9 ^{cd} _b	313 ± 21.9 ^{cd} _b	305 ± 21.2 ° _b	301 ± 22.6° _b	297 ± 26.2 ° _b
XSEA	400	96 ± 2.73° _a	353 ± 10.6 ^d _d	341 ± 10.6 ^d _{cd}	316 ± 7.78 ^d _b	291 ± 5.66 ° _b	267 ± 23.3 ^b _b	250 ± 23.3 ^b _b	220 ± 25.5 ^b _b
XSETH	400	99 ± 4.95° a	327 ± 9.2 ^d _c	320 ± 7.1 ^d _{bc}	312 ± 4.95 ^d _c	300 ± 7.1 ^d _b	288 ± 10.6° _b	278 ± 13.4 ° _b	266 ± 15.6° _b
XSW	400	112 ± 4.24° a	316 ± 8.5 ^d _d	314 ± 9.90 ^d _d	315 ± 12.73 ^d _d	310 ± 14.1 ° _b	309 ± 14.4 ° _b	308 ± 14.1° _c	306 ± 14.1 ° _c

The values are mean \pm SD, of triplicate experiments, the different letters over each range denote variation ($p \le 0.05$) in blood glucose level at various intervals (superscripted letters) and type of fraction administered (subscript letters).

XSETH, and XSW were measured as 325 \pm 21.2 mg/dL, 353 \pm 10.6 mg/dL, 327 \pm 9.2 mg/dL, and 316 \pm 8.5 mg/dL, respectively. There was no substantial decrease in the blood glucose level of the XSPE group (treated with petroleum ether) observed and two mice out of six died on days 8 and 13 of the treatment, while on day 18 the mean blood glucose level of the remaining mice decreased upto 297 \pm 26.2 mg/dL (9%). Animals treated with ethyl acetate (XSEA) showed a substantial decrease in the blood glucose level from day 1 353 \pm 10.6 mg/dL to 220 \pm 25.5 mg/dL (38%) on day 18. The blood glucose levels of animals treated with ethanol (XSETH) and water (XSW) fractions on the day 18 were 266 \pm 15.6 mg/dL and 306 \pm 14.1 mg/dL respectively.

Ethanol fraction lowers the blood glucose level by up to 19% while water fraction decreases it by only 3%. Overall, it can be claimed that the ethyl acetate fraction of XSL (XSEA) contains substantial amounts of phenolic bioactive antioxidants which decreased the blood glucose level of diabetic mice. However, more comprehensive studies with a larger number of mice might be helpful to generalize these findings.

In the previous studies, the aqueous leaf extract of XSL was also used as an antidiabetic agent at a dose of 500 mg/kg and 250 mg/kg body weight respectively in alloxan-induced diabetic mice for 10 days and a significant reduction in blood glucose level was observed at the

high dose of XSL aqueous extract (Mouhamad, 2022). Similarly, methanolic extract of XSL leaves prepared by the soxhlet apparatus was used to determine the antioxidant potential (in-vitro) and antidiabetic potential (in-vivo) in albino rats by oral administration at a dose of 200 mg/kg and 400 mg/kg body weight for 9 days (Umer et al., 2016). In another study hypoglycemic effect of ethanolic extract of XSL and its isolated compound were evaluated in alloxan-induced diabetic mice for 14 days and compared with Glibenclamide 10 mg/kg b. w; an antidiabetic drug (Ahmad et al., 2016). Leaf and fruit methanolic extract of X. strumarium obtained by the soxhlet method was used to measure the antidiabetic effect in diabetic mice by oral administration of 400 mg/kg b. w of extract for a period of 24 days (Harikumar et al., 2012). Methanolic extract of XSL stem was used to evaluate the hypoglycemic effect at doses of 100 mg/kg and 200 mg/kg body weight and antidiabetic potential was compared with Glibenclamide 0.6 mg/kg in Streptazocin-induced diabetic rats (Narendiran et al., 2011). By surveying the literature, it can be observed that there is no research work published about crude foliage fractionation and comparison of their antioxidant, antidiabetic potential, and variation in body weight. In the present research work, four fractions of XSL were obtained and their total phenolic content (TPC), antioxidant, and antidiabetic potential were studied. The antidiabetic potential of ethyl acetate fraction (XSEA)

fractions was comparable with standard Glibenclamide. By comparing results observed by all fractions, it has been observed that the XSEA fraction showed high TPC value, antioxidant potential and antidiabetic effect followed by the ethanol, petroleum ether, and aqueous fractions.

4 Conclusion

The aim of the current research was to assess the antidiabetic potential of different fractions of XSL foliage on diabetic mice. After 18 days of constant administration of XSL fractions, the blood glucose level of mice significantly decreased. These verdicts reinforced the traditional usage of the XSL foliage as an antidiabetic mediator as well as for the treatment of a number of ailments (Rahman et al., 2022). In order to boost insulin release from pancreatic cells, the conventional medication Glibenclamide has been used to treat diabetes for several decades. Several plant extracts were reported to exhibit hypoglycemic effects through stimulatory actions on insulin release, which is a possible pathway by which plant extracts lower blood glucose levels. This mechanism involves elevating the pancreatic insulin secretion from islets of Langerhans β cells or through its breakdown from the confined form (IfedibaluChukwu et al., 2020). According to some reports, other plants may also influence blood glucose levels by stimulating insulin release. Some wild herbs have chemical components that have hypoglycemic effects. There is evidence that certain types of plantderived compounds, including carboxyatractyloside, chlorogenic acid, caffeic acid, and other phenolic compounds lower blood glucose levels. It has been found that plant's extracts and metabolites have pharmacological effects. The results obtained supported the use of XSL in conventional medical system to cure diabetes. To determine the precise pathway of the anti-diabetic activity of the XSL fractions, a more comprehensive pharmacological and chemical research with large number of subjects is required.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary material, further inquiries can be directed to the corresponding authors.

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Ethics statement

The animal study was approved by Institute of industrial biotechnology Government College University, Lahore. The study was conducted in accordance with the local legislation and institutional requirements.

Author contributions

AS conducted experiments and prepared the draft, MM conceptualized and supervised this research, and SS, SA, AR, and AA reviewed the draft. All the authors endorsed the final version of the article.

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Conflict of interest

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Step-by-step guide for electrochemical generation of highly oxidizing reactive species on BDD for beginners

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Selecting the ideal anodic potential conditions and corresponding limiting current density to generate reactive oxygen species, especially the hydroxyl radical (*OH), becomes a major challenge when venturing into advanced electrochemical oxidation processes. In this work, a step-by-step guide for the electrochemical generation of *OH on boron-doped diamond (BDD) for beginners is shown, in which the following steps are discussed: i) BDD activation (assuming it is new), ii) the electrochemical response of BDD (in electrolyte and ferri/ferro-cyanide), iii) Tafel plots using sampled current voltammetry to evaluate the overpotential region where *OH is mainly generated, iv) a study of radical entrapment in the overpotential region where *OH generation is predominant according to the Tafel plots, and v) finally, the previously found ideal conditions are applied in the electrochemical degradation of amoxicillin, and the instantaneous current efficiency and relative cost of the process are reported.

KEYWORDS

electrochemical oxidation, highly oxidizing reactive species, sampled current voltammetry, boron-doped diamond, Tafel plot, amoxicillin degradation

1 Introduction

When delving into the wide world of advanced oxidation processes (AOPs), we found an overwhelming amount of information that can be frustrating for a beginner. All these technologies mainly seek the generation of the hydroxyl radical (*OH) and other highly oxidizing reactive species (HORS) to carry out highly efficient oxidation processes. Advanced electrochemical oxidation (AEO) is one of the most studied AOPs for the oxidation/degradation of organic matter in aqueous media (Martínez-Huitle et al., 2015; Bessegato et al., 2021; Hand and Cusick, 2021; Pierpaoli et al., 2021; Espinoza-Montero et al., 2023). AEO consists of applying a polarization potential to a conductive substrate (metal or semiconductor) to produce HORS, mainly *OH if possible, on the electrode-solution interface (Ganiyu et al., 2021; Lee et al., 2022; Xu et al., 2023b). One of the most appreciated electrodes for AEO is the boron-doped diamond (BDD) electrode. BDD is a p-type semiconductor due to the chemical nature of boron, as they are impurity/dopant donors, so in thermal equilibrium, there is an imbalance of charge carriers in the valence and conduction bands, with the positive holes being the majority carriers (Yokoya et al., 2005; Yang et al., 2019). As an electrode, BDD is a very versatile material as it has unique electronic properties. It is a non-active electrode; i.e., its chemical structure is not compromised by the

redox processes to which it may be subjected, although its reticular terminations are susceptible to slight changes during oxidation and reduction (Berenguer et al., 2019). As it has a very wide potential, this electrode has been used for multiple applications in electroanalytical chemistry (Wei et al., 2011b; Joshi et al., 2022), electrosynthesis (Ivandini and Einaga, 2017; Lips and Waldvogel, 2019), and AEO for water treatment (Cisneros-León et al., 2023).

For water treatment, electrochemical oxidation can take advantage of the complexity of mixtures of organic and inorganic species that can be found in typical contaminated water since, depending on the composition of the inorganic electrolyte in the system to be treated, the so-called HORS can be generated (Wen et al., 2017; Xu et al., 2021; Ahmed et al., 2022; Parvulescu et al., 2022; dos Santos et al., 2023; Mousset, 2023). Most HORS are "free" radicals and detrimental to most biological systems; the opposite is true for advanced oxidation processes, where they contribute to the oxidation of organic and inorganic matter (Murphy et al., 2022; Cisneros-León et al., 2023). HORS covers a wide spectrum of inorganic radicals, ions, and neutral molecules in aqueous media whose redox potentials are varied (Ross and Neta, 1982; Armstrong et al., 2015). Directly or indirectly, electrochemical oxidation can generate reactive oxygen species (ROS) such as the superoxide radical (O2 •-), singlet oxygen (1O2), hydrogen peroxide (H2O2), and the strongest oxidant, the hydroxyl radical (*OH) (Xie et al., 2022). On the other hand, in the presence of nitrates and nitrites, through the reaction of the electrogenerated OH, reactive nitrogen species (RNS) such as mono-, di-, and trioxide nitrogen radicals (NO[•], NO₂[•], and NO₃[•]) can be obtained (Wu et al., 2020; Rayaroth et al., 2022). Likewise, in the presence of sulfates, sulfites, and carbonates, sulfate (SO₃•-) and sulfite (SO₃•-) radicals, the persulfate anion (S2O82-), and the carbonate radical (CO30) can be generated in situ, and in the presence of chlorides or chlorites, reactive chlorine species (RCS) are very common, such as the chlorine radical (Cl°) and the oxychloride radical (Cl°) (Ganiyu and Martínez-Huitle, 2019; Ganiyu et al., 2021; Zhou and Xiao, 2022). In addition, reactive phosphate species (RPS) are generated indirectly by reacting OH with phosphates, producing phosphate radicals in three acidic forms, the H₂PO₄ radical being the most powerful oxidizing species of this type (Weiss et al., 2008). All these species can contribute greatly to the oxidation of organic and inorganic pollutants, achieving a predominant synergistic effect, or they can have the opposite effect since they all compete in coupled reactions in electrocatalysis.

This work intended to provide a step-by-step guide aimed primarily at beginners in the investigation of electrode properties in common electrolytic systems, with emphasis on the HORS that can be generated in the medium and that compete in catalysis. First, an approach from the existing theory in this field is addressed, and then, a real example is given using the BDD electrode in a known electrolytic environment.

2 Step-by-step electrooxidation on BDD

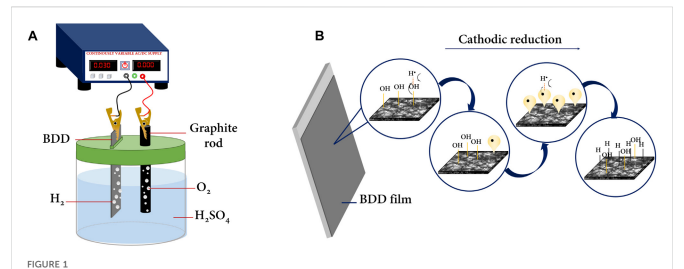
To experimentally determine the optimal operating conditions at AEO, some fundamental electrochemical tests in known redox environments must be considered. In the following, we will carefully

describe the suggested minimum steps that will allow us to develop our understanding of the optimal conditions for electrolysis at larger scales.

2.1 Electrode activation and cleaning (if required)

BDD is a "non-active" electrode when acting as an anode, and the HORS that are generated when a high oxidation overpotential is applied do not drastically change their chemical nature; this property makes it a unique material and is the basis for a large number of applications including in the area of environmental remediation (Sánchez-Montes et al., 2020; Carrera-Cevallos et al., 2021; Cisneros-León et al., 2023; Long et al., 2023). Therefore, studying the surface terminations of the crystal lattice of this material becomes key and is addressed in this section.

Diamond is a purely insulating material since its crystal lattice is composed of hybridized C-sp³. However, by introducing impurities of boron or nitrogen atoms during its synthesis using controlled methods at high temperatures and pressure, such as chemical vapor deposition (CVD) (Macpherson, 2015; Sun et al., 2021), a conducting, semiconducting, or superconducting material is obtained as a result of two factors; the first is that by introducing boron (or nitrogen) atoms, carriers that previously formed bonds are "released" and move freely through the lattice and participate in conduction in the BDD (Yokoya et al., 2005; Yang et al., 2019). On the other hand, when impurities are introduced to the diamond, it changes the hybridization of certain carbon atoms (from C-sp³ to Csp²), which extends to the surface terminations of the crystal lattice. The sp2 (C-sp2) hybridization (graphitic carbon) is the one that contributes the most to the conductive properties of BDD. Therefore, the surface termination of the BDD is a mixture of C-sp³ and C-sp². When the surface termination is predominantly C-sp³, the BDD is said to have an H-termination, while when the surface termination is C-sp², it is said that the surface termination is predominantly oxygen or oxygenated organic functional groups (Medeiros De Araújo et al., 2014; Garcia-Segura et al., 2015). Surface modifications of BDD from C-sp2 to C-sp3 or vice versa are possible through different controlled experimental methods. BDD electrode manufacturers often prefer methods that are as effective as their cost, e.g., oxygen plasma for oxygenated surfaces and hydrogen plasma for hydrogenated surfaces, but these are not practical for use in most laboratories (Kasahara et al., 2017; Einaga, 2022). On the other hand, surface modification of BDD electrochemically is simpler and cheaper; e.g., anodic polarization generates an oxygen surface termination, and cathodic polarization generates a hydrogen surface termination, for which a power source and a two-electrode cell are required (Figure 1A). This simple assay is commonly performed in strong acid media (H2SO4, HClO4, and HNO₃) at moderate concentrations. The transitions from H to O or vice versa, in addition to the intrinsic equilibrium of adsorption-desorption of the gases produced on the surface—O2 at the anode and H2 at the cathode—are achieved as long as the redox potentials of the organic forms present in the lattice are reached; i.e., care must be taken to control the DC density applied to the cell as well as the time for which the electrode is exposed to oxidation or reduction.



(A) Two-electrode cell for electrolysis in an acidic medium performed in lab. (B) BDD surface transition by cathodic reduction, adapted from the work of Einaga (2022).

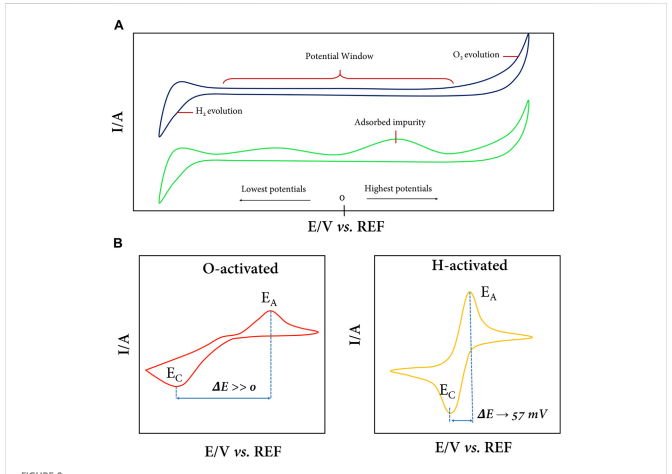


FIGURE 2
(A) Voltammetric profiles of the full potential window of the BDD in the presence of an electrolyte. Without the presence of contaminants or adsorbed impurities, only the oxygen and hydrogen evolution peaks are observed (top), while on a soiled surface, peaks are present in the capacitive zone (bottom). (B) Typical BDD CVs in the ferri/ferro-cyanide redox couple for an oxygenated (left) and hydrogenated (right) surface.

Einaga (2022) proposed a mechanism for the transition from oxygenated to hydrogenated BDD terminations. In essence, there is a homolytic contribution of hydrogen and the -O-H group from the BDD surface, yielding a good leaving group (water). In this way, the carbon formerly bonded to the R-CH₂-O-H group leaves a hybrid orbital with an unpaired electron ripe to overlap with another atomic hydrogen and obtain surface groups with hydrogenbonded carbons (R-CH₂-H) (Figure 1B). In addition, electrolysis in a strong acidic medium serves to clean impurities from BDDs prior to use, if required. It is very common to adsorb carbon impurities generated during the synthesis of the BDD or by the adsorption of contaminants from the environment or from contact with other surfaces. Therefore, it is common that during the manufacture of BDDs, especially bipolar (double-sided modified) BDDs, a thin layer of hybrid material with major impurities remains, and electrochemical cleaning is useful to remove this layer, as recommended by the manufacturers themselves. Cleaning, which should be understood interchangeably as surface activation of the BDD, is a soft electrochemical method that does not compromise the crystalline integrity of the BDD, much less of the substrate on which the doped diamond layer has been deposited, which is commonly Nb or n-type Si.

2.2 Electrode characterization

By cyclic voltammetry (CV), the cleanliness and surface termination of the BDD can be evaluated, as shown in Figure 2A. The BDD has an impurity-free surface (Figure 2A top) when no peak appears in the potential range of the capacitive region, and only the extreme peaks of oxidation evolution (more positive potentials) and hydrogen evolution (more negative potentials) should be detected. However, additional signals characteristic of surface carbon impurities often appear when the electrode is not properly cleaned/activated (Figure 2A bottom). These CV profiles are typically recorded in the presence of the electrolyte only, commonly Na₂SO₄, H₂SO₄, and phosphate buffer solutions, in dilute concentration (0.1 M). Although it is also possible to use strong bases (NaOH or KOH) as the supporting electrolyte to generate these voltammograms, caution must be exercised when using a suitable reference electrode (Zoski, 2007).

CV, in addition to being useful for calculating the electrode area by applying the Randles-Sevcik equation, is also useful for evaluating the predominant surface termination of H or O on BDD after electrochemical treatment. For this purpose, one can use the $[Fe(CN)_6]^{3-}/[Fe(CN)_6]^{4-}$ redox couple (or others) in dilute concentration and in a suitable electrolyte medium, typically KCl in a 1:100 ratio, although it is common to use more concentrated electrolyte solutions to avoid potential losses due to Ohmic drop (Suffredini et al., 2004; Actis et al., 2008; Zanin et al., 2014; Cheah and Chernev, 2021; Bard et al., 2022; Zheng, 2023). Figure 2B shows the typical CVs of BDD electrodes in the ferro/ferri-cyanide redox couple. At an electrode with oxygen surface terminations (Figure 2B left), the $[Fe(CN)_6]^{3-}/[Fe(CN)_6]^{4-}$ redox process becomes very slow, and the separation of anodic potential (Ea) and cathodic potential (E_c) peaks is enormous; i.e., ΔE_p (E_c-E_a) takes large values. On the contrary, in Figure 2B (right), a typical CV of a BDD electrode with H-terminations predominant on the surface is observed, where the redox couple response is fast, chemically and electrochemically reversible, $\Delta E_{\rm p}$ is close to 57 mV, and the width at half on the forward scan of the peak is close to 59 mV (Elgrishi et al., 2018). In practice, it is crucial to define which surface is of interest depending on the intended application of the BDD electrode (Oliveira et al., 2010; Kondo, 2022). For example, if a semiconductor-photocatalyst-modified electrode is required, it is ideal to have an oxidized surface as the physical and chemical interactions are stronger for oxygen with the semiconductor metal. On the other hand, if bare electrode electrocatalysis is what is sought, a predominantly hydrogenated surface is ideal since charge transfer at the solid–liquid interface is favored

2.3 Selecting parameters

Once BDD is electrochemically characterized to "micro-scale," the next step is the search for the "ideal" parameters to carry out electrochemical oxidation with the greatest efficiency, driven mainly by HORS, which will be discussed in depth later. AEO is a complex process involving the balance of several variables that must be carefully controlled in practice (Ganiyu et al., 2021; Bany Abdelnabi et al., 2022; Lee et al., 2022; Xu et al., 2023a; Zheng, 2023). In this section, we will focus mainly on the potential and/or current density to be applied to the system and how the electrochemical test should be carried out to find it. Our purpose is to systematically relate the Tafel analysis to the different electrolyte species that may be present and their consequent formation of HORS at the BDD electrode. For the construction of the Tafel curves, we will use voltammograms generated by sampled current voltammetry, a very effective and easy-to-apply technique.

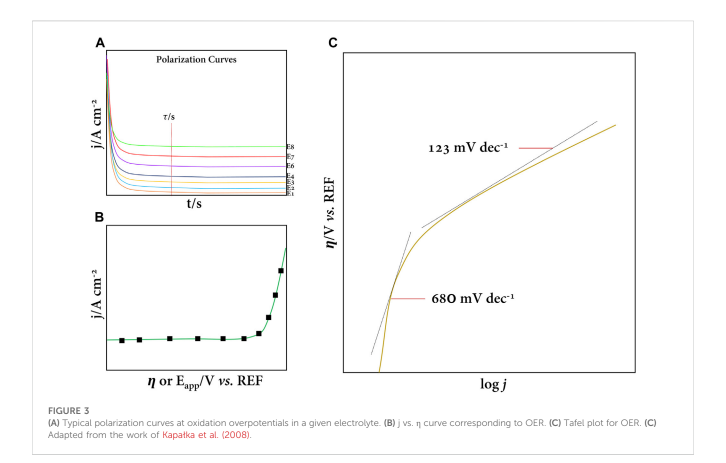
2.3.1 Tafel analysis

To monitor an electrochemical process quantitatively and from the point of view of kinetics, the Tafel analysis is often very useful for certain processes (Bard et al., 2022; Shih et al., 2022). The equation that quantitatively relates the potential to the current of a specific system is called the Tafel equation (Eq. 1), and it was first designed to study the hydrogen evolution reaction (HER) (Tafel, 1905; Petrii et al., 2007; Fang and Liu, 2014; Jung et al., 2022).

$$\eta = a - b \ln j \quad or \quad \eta = \frac{RT}{\alpha F} \ln i_0 - \frac{RT}{\alpha F} \ln j,$$
(1)

where η is the overpotential, which is calculated from the algebraic subtraction of the potential applied (E_{app}) to the working electrode and the open-circuit potential (E_{OCP}) , *i.e.*, the equilibrium potential that is specific for each electrolyte in solution. The Tafel plot is obtained by plotting $\ln j$ vs. η , where b is the Tafel slope(s), which is of interest to us because of its kinetic significance (Fang and Liu, 2014).

To study the kinetics of redox processes in electrochemistry, the rotating disk electrode is employed since the speed of revolution is controlled and the diffusional and capacitive influences of the system are eliminated, thus allowing the kinetic parameters and charge transfer constants to be calculated with high reproducibility (Chen et al., 2020). However, not all laboratories have this



sophisticated system at their disposal. Instead, sampled current voltammetry (SCV) is a useful technique that frees us from this experimental drawback, and the kinetics of reversible or irreversible reactions can be studied with the Tafel analysis without major problems (Soares et al., 2020; Rodríguez and Denuault, 2021).

An SCV assay consists of subjecting the working electrode to various polarization potentials—in a specific range and in the medium of interest—during a specific pulse time (chronoamperometry); in this sense, the so-called polarization curves are obtained (Figure 3A). The polarization curves are used to select the current density corresponding to each potential at a given time constant (τ) , thus plotting the η vs. j or E_{app} vs. j curve (Figure 3B) and applying the Tafel analysis to the data obtained.

Kapałka et al. (2008) pioneered the study of the kinetics of the oxygen evolution reaction (OER) at a BDD electrode (Figure 3C). They showed that in the Tafel analysis, there were two well-defined slopes corresponding to independent, one-to-one electron transfer processes. For our interest and depending on the electrolyte, the Tafel analysis gives us a hint of the optimal potential to produce HORS before oxygen evolution is favored (thermodynamically). Therefore, this kinetic study will allow us to select a wide range of anodic potentials and evaluate the production of HORS, as discussed below.

2.3.2 HORS at BDD

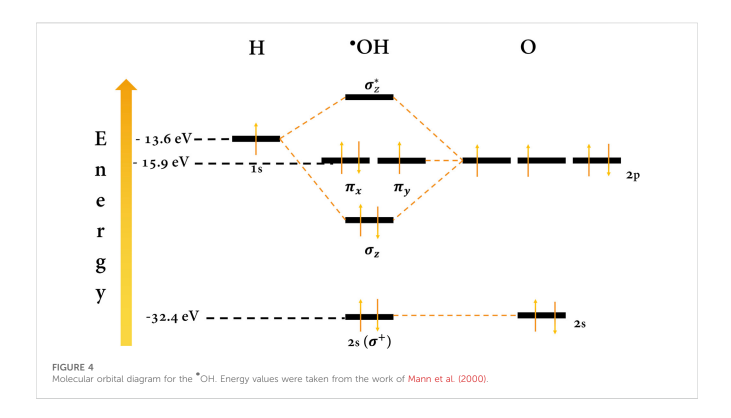
A further digression is necessary here to address the HORS generated on the BDD. Depending on the electrolyte in which the electrolysis is carried out, different reactive species capable of efficiently oxidizing organic molecules may coexist, some more

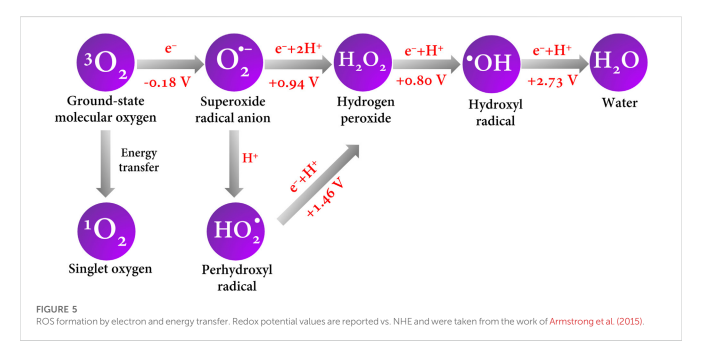
than others. We cannot begin this discussion without first analyzing the ROS and, in essence, the hydroxyl radical (*OH). This radical is the first intermediate of the OER from water splitting (Reaction 1), and it is generated in a process favored by kinetics rather than thermodynamics; i.e., large overpotentials are required for its formation on the BDD surface (Siahrostami et al., 2017; Zhu, 2019; Xie et al., 2022).

$$BDD + H_2O + E_{app} \rightarrow BDD(OH^*) + H^+ + 1e^-$$
 (1)

BDD (OH*) refers to the OH adsorbed on the surface of the BDD, specifically on the active site, which plays a crucial role (Moreira et al., 2017; Vogt and Weckhuysen, 2022). OH is a highly reactive and unstable species whose lifetime is estimated to be a few nanoseconds (Moreira et al., 2017). The high reactivity of the OH is explained by its unpaired electron in an antibonding orbital (π^*) of an oxygen atom located at high energy (Figure 4). This instability (in energy) means that its electron easily pairs with an organic species to form covalent bonds in the electrooxidation pathway (Imlay, 2003; Krumova and Cosa, 2016). Its oxidizing power (~2.73 V vs. NHE), second only to fluorine, is due to its high tendency to capture electrons and stabilize the antibonding orbital. Therefore, it is capable of non-selectively oxidizing a wide variety of organic pollutants as long as the redox potential of the target molecules is aligned with the redox potential of the radical at the interface (Armstrong et al., 2015).

On the other hand, molecular oxygen in its fundamental state is triplet $(^3O_2)$ and is dissolved in the medium or generated by the electrolysis of water; it can trigger reduction reactions to generate other highly reactive species but with lower oxidizing power than





the *OH (Krumova and Cosa, 2016). When O_2 is reduced by the conduction band electrons of the BDD, on transfer of a first electron, it produces the superoxide anion radical $O_2^{\bullet-}$ (E⁰ = -0.18 V vs. NHE) (Armstrong et al., 2015; Hayyan et al., 2016). In brief, Figure 5 shows a schematic of the ROS generated by electron transfer and their redox potentials.

In short, the *OH, due to its oxidizing power, is the species that is sought to be generated efficiently in AEO, and it will be seen from now on that this radical plays a crucial role in the generation of other highly reactive oxidizing species (Gligorovski et al., 2015). As for pH, it is also a

parameter that greatly influences the generation of reactive species and electrocatalysis in general. We said that the ${}^{\bullet}$ OH production process is kinetically favorable, where pH influences the rate of charge transfer (Tao et al., 2019; Zhang et al., 2020). At low pH, the concentration of ${}^{\bullet}$ OH. † O ${}^{+}$ increases, which favors the generation of ${}^{\bullet}$ OH. † O ${}^{+}$ can react with electrons from the electrode (depending on the electrode potential) and produce atomic hydrogen, which is highly reactive and reacts with water to produce ${}^{\bullet}$ OH (Katsounaros et al., 2014). Very low pH can also be detrimental to the electrode as it can create a passivation layer on the electrode surface that limits charge transfer; in BDD, this is not a

drawback as its high stability in strong acid environments frees it from this issue (Xie et al., 2022). Other considerations must be taken with respect to pH since the stability of the target molecule to be degraded is also a tremendously influential factor, as well as the concentration of the electrolyte species in the solution (Fornaciari et al., 2022).

The utility of ions in a solution is imperative for the vast majority of electrochemical processes (Bard et al., 2022). In the process at stake here, the supporting electrolyte plays a dual role since it is not only responsible for mitigating the migration of the electroactive species and serving as an ionic conductor in the medium but the ions can also react instantaneously with the electrogenerated OH and produce other chemical species of high reactivity and, sometimes, high oxidizing power (Kim et al., 2018; Hand and Cusick, 2021). Many authors refer to this process as the "activation" of a certain species since in the absence of any perturbation, the ions are not a major threat to organic pollutants (Lin and Deng, 2021). In a medium containing sulfate ions, activation of the sulfate radical (SO₄[•]) and other reactive sulfate species is very likely. This radical anion is mainly produced by the oxidation of (SO_4^{2-}) in the presence of ${}^{\bullet}OH$ adsorbed on the BDD in a 1-electron transfer process (Reaction 2). Another species, the persulfate anion $(S_2O_8^{2-})$, can be generated by direct oxidation on the surface of the BDD or by oxidation of sulfate by the OH in a 2electron transfer process (Reaction 3) or by direct combination of two sulfate radicals (Reaction 4) (Oh et al., 2016; Lee et al., 2020; Ganiyu et al., 2021; Araújo et al., 2022).

$$BDD(OH^*) + SO_4^{2-} \rightarrow BDD(SO_4^{\bullet-}) + OH^{-}$$
 (2)

$$2BDD(OH^*) + 2SO_4^{2-} \rightarrow BDD(S_2O_8^{2-}) + 2H_2O$$
 (3)

$$2BDD(SO_4^{\bullet-}) \to BDD(S_2O_8^{2-}) \tag{4}$$

In the literature, the vast majority of works that seek to treat polluted water by electrochemical oxidation use sulfate-based electrolytes, typically sodium or potassium sulfate, or sulfuric acid. Why is this? This can be accounted for by the oxidizing power of this radical, which ranges from 2.5 to 3.1 V vs. NHE; these values compete with the oxidizing power of the OH (Ganiyu and Martínez-Huitle, 2019; Ganiyu et al., 2021; Araújo et al., 2022). When comparing these two species, both radicals have an affinity for capturing electrons in redox processes since they have unpaired antibonding molecular orbitals in oxygen, while the electronegativity of S is greater than that of O, and the polarization of the bond is more efficient (Lin and Deng, 2021). Therefore, if these two species can coexist in a moderate "balance," the treatment of contaminated water can be very efficient. Understanding the oxidation power of these species from theory can be a simple issue, but, in practice, it is not so since variables such as pH, applied potential and current density, temperature, and contaminants must be controlled.

Reactive chlorine species (RCS) are produced in an electrolytic chloride environment. Mainly, the chlorine radical (Cl[®]) is the outstanding species of this group with an oxidation power of approximately 2.43 V vs. NHE (Armstrong et al., 2015). It is produced by indirect oxidation, *i.e.*, by the oxidation of Cl⁻at the electrode surface by the adsorbed [®]OH (Reaction 5) (Mostafa et al., 2018). On the other hand, the hypochlorite radical (ClO[®]), also a highly reactive species of chlorine, is produced by the same oxidation reaction by the [®]OH (Reaction 6) and has a relatively low oxidation power (1.39 V vs. NHE) (Armstrong et al., 2015). The [®]OH can trigger reactions with chloride in the medium to produce hypochlorite,

chlorite, chlorate, and perchlorate anions, but they are not highlighted here since their oxidizing power is overshadowed by their toxicity, and reactions producing these species can be found anywhere (Ganiyu et al., 2021; Choo et al., 2022; Barnum and Coates, 2023).

$$BDD(OH^*) + Cl^- \rightarrow BDD(Cl^{\bullet}) + OH^-$$
 (5)

$$BDD(OH^*) + ClO^- \rightarrow BDD(ClO^{\bullet}) + OH^-$$
 (6)

Similarly, reactive phosphate species (RPS) are produced by hydroxyl radical chain reactions (Reactions 7–9). The phosphate radical (PO $_4^{\bullet 2-}$) can exist in its three acidic forms, with the radical (H $_2$ PO $_4^{\bullet}$) having the highest oxidizing power, approximately 2.75 V vs. NHE (Armstrong et al., 2015). The other radical species are less reactive for both redox processes and acidic proton capture (Ross and Neta, 1982; Weiss et al., 2008).

$$BDD(OH^*) + H_3PO_4 \rightleftharpoons BDD(H_2PO_4^{\bullet}) + H_2O$$
 (7)

$$BDD(OH^*)H_2PO_4^- \rightleftarrows BDD(H_2PO_4^{\bullet}/H_2PO_4^{\bullet-}) + OH^-/H_2O \quad (8)$$

$$BDD(OH^*) + HPO_4^{2-} \rightleftharpoons BDD(HPO_4^{\bullet-}/PO_4^{\bullet2-}) + OH^-/H_2O \quad (9)$$

Finally, it is not difficult to imagine that reactive nitrogen and carbon species are produced by the indirect oxidation of the hydroxyl or sulfate radical with nitrates, nitrites, and carbonates to generate nitrogen di- and trioxide radicals (NO_2^{\bullet} , NO_3^{\bullet}), as well as the carbonate radical ($CO_3^{\bullet-}$) (Reactions 10–12). Among these species, the nitrogen trioxide radical is the strongest oxidant (2.5 V vs. NHE) (Armstrong et al., 2015). Despite being weaker than the hydroxyl radical, they can oxidize on one-electron transfer reactions a wide variety of organic and inorganic molecules, such as phenols, anilines, sulfur compounds, and some metal ions (Ross and Neta, 1982; Fang et al., 2014; Rayaroth et al., 2022).

$$BDD(OH^*) + NO_2^- \rightleftarrows BDD(NO_2^\bullet) + OH^-$$
 (10)

$$BDD(OH^*) + NO_3^- \rightleftarrows BDD(NO_3^\bullet) + OH^-$$
 (11)

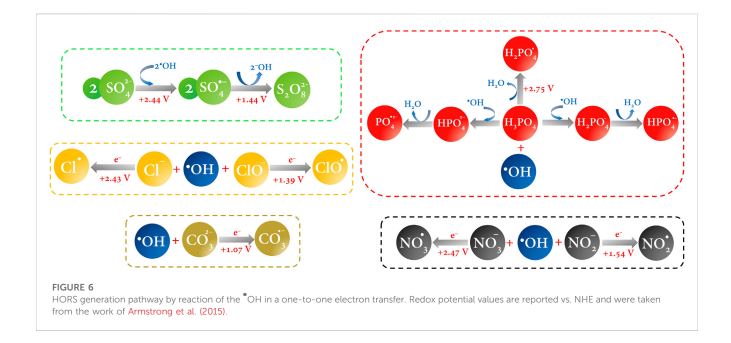
$$BDD(OH^*) + CO_3^{2-} \rightleftharpoons BDD(CO_3^{\bullet-}) + OH^{-}$$
 (12)

In this section, we left for a while the main aim of this work and tried not to digress in exposing all the species that can exist in an electrochemical reaction medium; instead, we emphasized the main highly reactive oxidizing species of each type depending on the electrolyte (Figure 6). In practice, it is obvious that after the ROS and, especially, OH, the other species are the consequence of the redox reaction of the latter with the ions in solution; *i.e.*, the other species can be regarded as trapping hydroxyl radicals. This is because OH is not selective and reacts simultaneously with whoever stabilizes its unpaired electron most quickly and efficiently. All these species highlighted above compete in electrocatalysis, and attributing the major contribution to pollutant degradation to just one of them would be a mistake. Instead, emphasis should be placed on controlling the other variables that take place in electrochemical oxidation.

3 Beyond the lab

3.1 Limiting current and limiting potential

All the previous analyses can be carried out successfully with accuracies that slightly deviate from the theory, but when the



results are transferred to a scale larger than that of the laboratory, it often involves complications that lead to results that deviate greatly from those predicted at the micro-scale, but are not wrong for the most part. These noticeable differences, which are often found at a real scale, are mainly due to the electrochemical cell and the intrinsic energy consumption. That is, when testing electrode behavior, electrolyte effects, and cell configuration in the laboratory, it is common to use simple systems on account of having known and small electrode areas, solution volumes no larger than 25 mL, and simple cell configurations, usually batch type (Bard et al., 2022). In the pre-pilot scale, all the dimensions of the upstream electrochemical components are increased. This increase has important implications for current and potential density. Thus, at scales larger than the laboratory, it is common to have control of the experimental limiting current, i.e., the current that maximizes the charge transfer efficiency of the redox reaction occurring at the electrode surface. Typically, the limiting current can be found in electrooxidation systems as a function of time in the chemical oxygen demand (COD) (Eq. 2) (Wei J. J. et al., 2011; Ochoa-Chavez et al., 2018).

$$i_{lim} = i_{lim}(t) = 4Fk_{m}COD(t), \qquad (2)$$

where i_{lim} is the limiting current, F is Faraday's constant, and k_m is the mass transfer coefficient. For microelectrode or rotating disk arrays, the limiting current can be calculated by SCV in known redox systems, and its procedure can be found elsewhere (Soares et al., 2020; Bard et al., 2022).

Then, when scaling a system, it is not strange that current densities or overpotentials reach values higher than those predicted by Tafel curves or voltammetric profiles in various electrolytes (Moreira et al., 2017). This is because the diffusion flow of the species within the solution towards the electrode surface changes and, therefore, the energy barrier to be overcome also increases; on the other hand, the potential loss by Ohmic

drop also increases significantly due to the resistance of the solution and the micro- and nano-bubbles that evolve on the surface, as well as the presence of the target molecule to be electrochemically oxidized (Petrii et al., 2007; Lee and Bazylak, 2021; Zheng, 2023). In practice and when disk electrodes are not available, the limiting potential and its limiting current are controlled in a non-analytical way (at least for now); *i.e.*, when the electrooxidation test is carried out, the electrode potential is progressively increased, and the current density circulating in the reactor is controlled, and when the proportionality between the potential and current is lost, the limiting current has been exceeded; in other words, the maximum potential value that the system supports so that the charge transfer is also maximum corresponds to the current limit where the process is optimized.

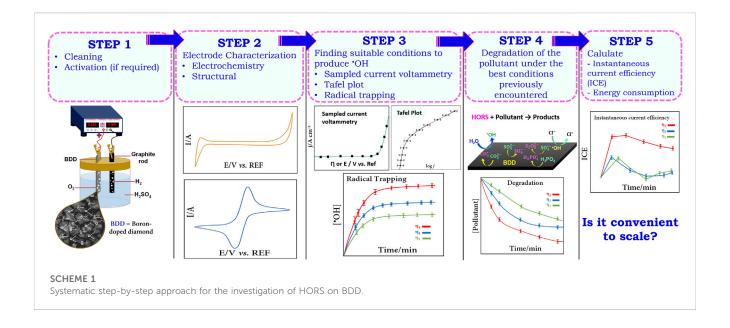
3.2 Instantaneous current efficiency and energy consumption

Additionally, another parameter of interest when scaling certain processes is the determination of the instantaneous current efficiency (ICE), which gives us information regarding the optimum range of applicability of a process (Comninellis and Pulgarin, 1991; Xing et al., 2018); this can be calculated by the following equation:

$$ICE = \frac{[COD_t - COD_{t+\Delta t}]FV}{8I\Delta t},$$
(3)

where $(COD)_t$ and $(COD)_{t+\Delta t}$ are the chemical oxygen demand at times t and t + Δt (in g O_2 dm⁻³), F is Faraday's constant, V is the volume of electrolyte (dm³), and I is the applied current (A).

Furthermore, the energy consumption of the process is also relevant since it gives us a picture of the efficiency from the perspective of energy expenditure and also helps us to optimize



not only the parameters but also the resources (Brillas et al., 2009). Energy consumption is calculated by the following equation:

Energy consumption (kWh m
$$^{-3}$$
) = $\frac{E_{cell}It}{V_s}$, (4)

where E_{cell} is the cell voltage (V), t is the electrolysis time (h), and I is the current passing through the cell (A). Therefore, the cost of the process can be obtained according to the following equation:

Cost of process = Energy consumption
$$\times$$
 Cost per kWh. (5)

In summary, Scheme 1 presents a systematic representation of the steps that we aimed to introduce so far and serves as a guide for the user; we emphasize that this is a preliminary approach to get to know our working electrode in the presence of a suitable electrolyte. We encourage the authors to include new electrochemical and non-electrochemical techniques to gain a deeper understanding of the electrode materials to be investigated.

The following is a specific case study involving BDD in the search for the optimal potential range for the generation of HORS, applying the reasoning that has been shown so far in the search for the prime parameters by carrying out electrochemical tests that can be handled without major problems in any electrochemical laboratory.

4 Case study: BDD in a Na₂SO₄ medium

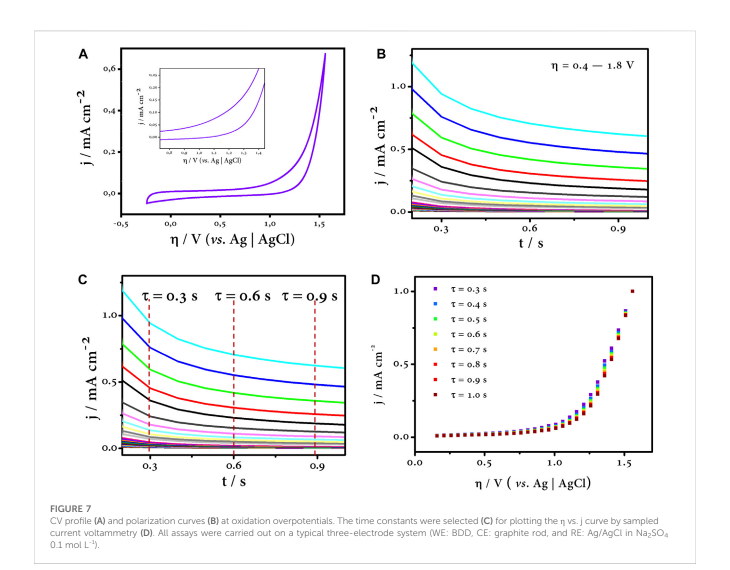
Quite recently, Lu et al. (2023) highlighted the most recent advances in the effects of electrolytes in a solution for electrochemical processes and discussed their effects on different reactions of interest in electrochemistry, such as oxygen and hydrogen evolution reactions (HER and OER, respectively) and oxygen and CO₂ reduction reactions (ORR and CO₂RR, respectively). Nevertheless, to date, no review has been found in the literature that addresses, in a simplistic fashion and with an experimental approach, the investigation of ionic species in a

solution and their participation in electrochemical oxidation as a guide for beginners. In this section, a real experimental example is presented as a case study that evaluates the behavior of BDD in Na_2SO_4 and other electrolytes of interest, emphasizing the steps that should be followed if one wishes to begin in this field.

Anhydrous sodium sulfate (N2SO4 99%) was purchased from J.T. Baker, N,N-Dimethyl-4-nitroso-aniline (for synthesis) was purchased from Merck, and amoxicillin was purchased from Sigma-Aldrich. All chemicals were used without further purification. BDD electrodes were purchased from CONDIAS and were cut into 25 mm × 15 mm rectangular plates, and an effective bipolar electrode area of 6.0 cm² in a cell volume of 60.0 mL was used for all experiments. The BDD was previously electrochemically activated by anodic polarization for 10 min in $0.5 \text{ mol L}^{-1} \text{ sulfuric acid } (\text{H}_2\text{SO}_4 95\%-97\%, \text{Merck}) \text{ at } 0.1 \text{ Acm}^{-2} \text{ in a}$ BK Precision AC/DC power supply. All electrochemical assays were carried out on a BioLogic potentiostat/galvanostat workstation in a typical three-electrode cell. The BDD served as the working electrode, Ag/AgCl served as the reference, and a graphite rod served as the counter electrode. For the degradation of amoxicillin (40 μM), a two-electrode system was used, with a BDD as the anode and a Pt mesh as the cathode.

4.1 Cyclic voltammetry and polarization curves

Once the electrode or electrode material to be evaluated—BDD in this case—has been selected, the electrolyte medium in which the behavior of the electrode is to be investigated is chosen without any additional species in the solution. Therefore, the first step is to perform cyclic voltammetry at oxidative potentials—actually overpotentials—in the chosen medium; our interest is in Na_2SO_4 for reasons already discussed earlier. The CV gives us a first insight into the potential where oxygen evolution is kinetically and thermodynamically predominant. The inflection point of the



oxidation branch of the CV indicates the potential where the transition from one electron transfer process to another is likely to occur and is likely to be the potential where most intermediate reactions compete, which is approximately 1.20 V vs. Ag/AgCl according to Figure 7A. Then, by chronoamperometry, we determine the polarization curves at the oxidation potentials determined in the previous CV profile (Figure 7B).

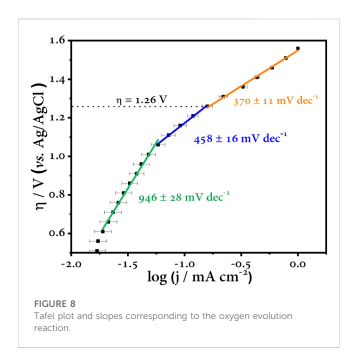
4.2 Sampled current voltammetry and $\boldsymbol{\eta}$ vs. j curves

After completing the polarization curves at the oxidation overpotentials, the next step is to choose a time interval, called time constant (τ) , in which the process is controlled by charge transfer (Bard et al., 2022). For this purpose, the SCV is a useful tool that allows us to control this phenomenon when a rotating disk electrode is not available (Soares et al., 2020). Typically, it consists of choosing current values at small time constants and constructing η vs. j curves (Figures 7B, C). In this range, the process is kinetic-controlled, and the influence of diffusion, *i.e.*, mass transfer, is limited, so the effect of the electrode capacitance does not affect our test, and all measurements are focused entirely on the Faradaic

current (Soares et al., 2020; Rodríguez and Denuault, 2021; Bard et al., 2022). Finally, Figure 7D shows all the η vs. j curves corresponding to different time constants, and all of them fit the same trend.

4.3 Tafel plot

As discussed in previous sections, the Tafel analysis is useful to evaluate the kinetics of an electrochemical process. In this sense, and summarized in the last step of our survey, the Tafel curve is constructed by plotting the logarithmic function of the current *versus* the overpotential (Figure 8). Three zones can be distinguished in this curve (highlighted by the straight lines). Each region of the curve has an associated slope, and this is the Tafel slope. The green line corresponding to the first zone has a slope value of 946 mV dec⁻¹, and this region is purely capacitive (it can be checked with the CV profile). The overpotential values up to about 1.1 V vs. Ag/AgCl are potentials where the double layer of the electrode—at the solid–liquid interface—is charged, and all this charge is stored in this imaginary region of the electrode, the inner Helmholtz plane (Bard et al., 2022). The next region illustrated by the blue line has a Tafel slope of 458 mV dec⁻¹, and this sharp change in slope indicates that another

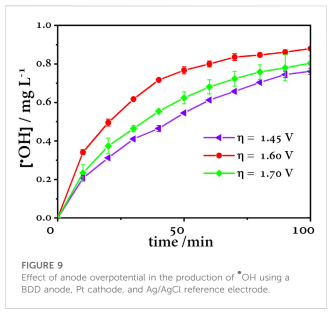


process(es) is taking place at the electrode surface. Finally, the last zone highlighted by the orange line has a slope of 370 mV dec⁻¹. This region plotted at overpotentials above 1.3 V vs. Ag/AgCl refers to the last process. This zone is distinctive as oxygen evolution reaction is predominant (Kapałka et al., 2008). This is self-evident as the error bars in this last region are almost undistinguishable, *i.e.*, the current and potential values do not oscillate to a large extent as long as the test is run again. The opposite is true for the previous zones. The intermediate zone is of interest to us as the former is easily discarded because of its capacitive contribution. The intermediate zone allows us to have an overview of the processes that can take place at these potential values. Since we know the chemical nature of the medium in which the test was carried out, we can say that this region is dominated by the generation of HORS that compete with the OH, our main oxidant.

However, despite our efforts, we cannot be certain of the exact potential at which the most highly oxidizing species can be harnessed for any given process, and to do so would be a tremendous mistake. Instead, these steps allow us to determine a wide range of potentials and discriminate those that do not contribute to our process to save us errors, time, and costs of unnecessary laboratory testing. Quantitative assays such as the quantification of reactive species or degradation of a known contaminant should be added to this procedure to extend this assumption and have a more adequate approach to our purpose; with this, we will have approached our search for the optimal experimental conditions to achieve adequate efficiencies in our advanced electrochemical oxidation process. To experimentally verify the overpotential region where the highest amount of *OH is generated, a radical trapping study was performed, as shown below.

4.4 Trapping of radicals

The *OH concentration was quantified by the N,N-Dimethyl-4-nitroso-aniline (RNO) method (Ochoa-Chavez et al., 2018). Anode overpotentials were chosen based on the Tafel plot at values above



1.26 V vs. Ag/AgCl, according to Figure 8. At an anodic overpotential value of 1.60 V vs. Ag/AgCl, the highest concentration of *OH is generated, while at 1.45 V vs. Ag/AgCl, the concentration is lower because the applied potential is not enough to generate the maximum concentration of *OH. On the other hand, at 1.70 V, the concentration decreases because OER is the predominant reaction (Figure 9).

4.5 Amoxicillin degradation

vIn order to verify that the higher *OH radical production matches a maximum degradation efficiency, the electrooxidation of amoxicillin (AMX 40 µM) was tested. The electrochemical degradation of amoxicillin was evaluated under the best OH production conditions, as discussed above. The degradation was followed by UV-Vis spectroscopy and chemical oxygen demand (COD). According to Figure 10A, at η = 1.40 V, the competition between the OH electrogeneration (E° = 2.43 V vs. SHE) and formation of sulfate species SO_4^{\bullet} (E° = 2.44 V vs. SHE) and $S_2O_8^{2-}$ (E° = 1.44 V vs. SHE) is expected to be carried out (Armstrong et al., 2015); this agrees with the first minutes of electrolysis where amoxicillin degradation decreases rapidly and then remains unchanged. This is because sulfate species are not able to completely mineralize highly stable small organic molecules such as formic acid and oxalic acid (k = 1.3×10^8 and 1.4×10⁶ L mol⁻¹ s⁻¹, respectively), reaching 24.1% AMX concentration removal and 55.8% COD removal in 360 min of electrolysis (Figure 10B). On the other hand, at $\eta = 1.60 \text{ V}$, the highest amoxicillin removal is achieved; this potential corresponds to the highest mineralization, and these removals decrease linearly until reaching 39.9% and 77.9%, respectively; therefore, this overpotential obtains the highest *OH production. Whereas, when overpotential increases, at $\eta = 1.80 \text{ V}$, the removal behavior is similar as that at 1.40 V, and in this case, the OER is predominant, obtaining only 36% amoxicillin removal and 54% COD removal.

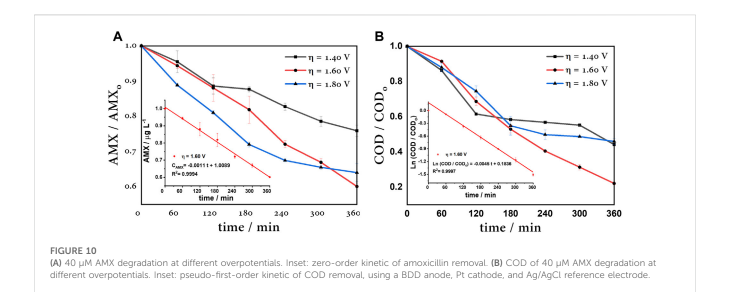


TABLE 1 Kinetic parameters of amoxicillin degradation.

η/V vs. Ag/AgCl	AMX remotion (%)	$k_{amx}/min \times 10^{-3}$	COD remotion (%)	$k_{COD}/min^{-1} \times 10^{-3}$
1.4	24.1	0.7	55.8	2.0
1.6	39.9	1.1	77.9	4.5
1.8	36.0	1.0	54.0	2.3

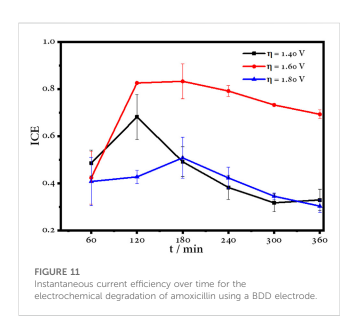
AMX, amoxicillin; η , overpotential; COD, chemical oxygen demand; k_{AMX} , zero-order kinetic constant; k_{COD} , pseudo-first-order kinetic constant.

In summary, Table 1 shows the kinetic parameters, where rate constants for amoxicillin degradation were adapted to a zero-order kinetic model and a pseudo-first-order kinetic model to COD removal. The COD constant rate of 1.60 V was almost two times greater than others' overpotentials; therefore, this is considered the optimal anode overpotential. In this case, this is the overpotential corresponding to the limit current, *i.e.*, the current associated to the maximum capacity to degrade AMX.

4.6 Instantaneous current efficiency and energy consumption

Figure 11 shows the evolution of the instantaneous current efficiency (ICE) with respect to electrolysis time, where maximum values of ICE are reached in the first hours, after which it decays due to the oxidation process and the formation of organic compounds. In the case of 1.60 V, it can be observed that after 2 h, the ICE remains stable, so the generation of *OH is continuous and allows progressive degradation, as it was discussed previously. Nevertheless, the ICE of the other overpotentials decreases drastically.

On the other hand, Table 2 shows the energy consumption and operating cost for the degradation of amoxicillin at different overpotentials. The cost of operation was calculated based on the cost per kilowatt-hour in Ecuador, which is 0.096 USD until March 2023 (Global Petrol Prices, 2023).



The results showed that the energy used during the electrolysis under the optimal overpotential was $0.224 \, \text{kWh m}^{-3}$, and the operation cost was $0.022 \, \text{USD m}^{-3}$, which represents the lowest cost among the other potentials tested and is in mutual agreement with the most efficient potential of the degradation process.

TABLE 2 Parameters of efficiency for the amoxicillin electrooxidation during a 6 h reaction.

ı	η/V vs. Ag/AgCl	ICE	Energy consumption/kWh m ⁻³	Cost/USD m ⁻³
	1.40	0.33	0.252	0.024
	1.60	0.69	0.224	0.022
	1.80	0.30	0.324	0.031

η, overpotential; ICE, instantaneous current efficiency.

5 Conclusion and outlook

Properly understanding the behavior of an electrode in a specific system can sometimes be a complicated endeavor, even more so when one does not know the charge transfer processes that may or may not take place in the medium under study and when this medium is a complicated mixture of organic and inorganic species. Electrochemical oxidation is a useful and efficient technology for many remediation applications if applied correctly, i.e., having adequate knowledge and control over the parameters that influence the process, both internal and external. In this sense, this work attempted to provide a step-bystep guide, particularly for beginners, on the tests to be considered in the investigation of electrodes and new electrode materials for advanced electrochemical oxidation processes, with emphasis on the range of anodic potentials where the generation of highly reactive oxidizing species predominates, which are priceless in these processes. It is not our intention that this guide should be followed rigorously, but rather that it should serve as a support for those who wish to get started in this field, adjusting these steps to their research objectives. The study of the kinetic behavior of BDD in different electrolytes by SCV, the Tafel plot, and trapping of radicals is essential to achieve results of this magnitude. A detailed study of the effect of the electrolyte is in progress in our research group.

Although this guide is particularly focused on the BDD, it is not limited to this electrode. This guide can be followed to investigate a new electrode material from an electrochemical point of view at first glance. In addition to the steps proposed here, the reader is free to implement other electrochemical techniques that suit his or her research. For example, electrochemical impedance spectroscopy is important when specific reaction mechanisms need to be studied and can provide valuable information about a particular electrode; however, this technique is beyond the scope of this work.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material; further inquiries can be directed to the corresponding author.

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Author contributions

GC-C: formal analysis, investigation, methodology, and writing-original draft. CP-C: data curation, formal analysis, investigation, methodology, and writing-original draft. EM: funding acquisition, investigation, validation, and writing-review and editing. PE-M: conceptualization, data curation, funding acquisition, investigation, methodology, project administration, resources, supervision, validation, writing-original draft, and writing-review and editing.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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