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Re-engineering luminol: new frontiers in chemiluminescence chemistry

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Luminol and its derivatives have emerged as powerful chemiluminescent agents with broad applications in biomedical diagnostics, forensic science, and environmental monitoring. Despite their widespread use, luminol's limitations, including poor solubility, short luminescence duration, and sensitivity to environmental conditions, have driven extensive research into the synthesis of more efficient derivatives. This concise review presents recent advances in the molecular engineering of luminol derivatives, focusing on design strategies that employ electronic modulation (e.g., introduction of electron-donating or withdrawing substituents) and steric tuning (e.g., alkylation and ring substitutions) to optimize its chemiluminescence efficiency, kinetics, emission wavelength, solubility, stability, and applicability for specific environments (e.g., biological systems). The review also discusses how these structural modifications impact luminol's performance within integrated systems, including forensics, bioimaging platforms, immunoassay technologies and microfluidic sensors, thereby linking molecular-level design with macroscopic function. Emerging macromolecular and polymer-based luminol systems, such as those incorporating hydrophilic carriers, nanoparticles, enzyme-responsive linkers, surface-immobilized polymer brushes, and multifunctional hybrid platforms, are also highlighted for their potential to overcome solubility and biocompatibility barriers while enabling targeted delivery or signal amplification. Finally, key challenges and future perspectives are outlined, including the development of near-infrared-emitting derivatives, improved storage stability, and interdisciplinary strategies for translating luminol chemistry into next-generation diagnostics and environmental sensing platforms. By summarizing these advancements, this review underscores the evolving role of luminol chemistry in modern analytical science and its potential to revolutionize next-generation detection technologies.

Design, System, Application

This concise review distils recent progress in the molecular design of luminol derivatives, with an emphasis on structural modifications that enhance chemiluminescence performance. By organizing current synthetic strategies and mechanistic insights into a coherent framework, the review offers a system-level understanding of how electronic and steric modifications influence luminol's luminescence properties. It also explores emerging applications in diagnostics, sensing, and environmental monitoring, and highlights future directions such as near-infrared emitters and lab-on-a-chip integration. As such, the review serves as a valuable design resource for researchers developing soft molecular systems with tuneable optical responses for healthcare and environmental applications.

1. Introduction

Chemiluminescence (CL) has recently emerged as an advanced strategy for detecting various analytes, including heavy metals, pollutants, metabolites, and pharmaceuticals. It has become a standard tool in environmental analysis,¹ forensic science,² clinical diagnostics and theranostics,^{3,4} as well as pharmaceutical and biomedical research.^{5,6} This widespread application is attributed to its simplicity,

background-free nature, ultra-high sensitivity (down to attomolar concentrations), rapid detection, and flexible integration into diverse analytical strategies.^{5,7,8}

CL detection relies on a chemical reaction that generates electronically excited species, which emit light upon returning to their ground state. Luminol (3-aminophthalhydrazide) is the most extensively chemiluminescent reagent due to its broad range of applications. 4,9 Its oxidation is triggered by specific reactive oxygen and nitrogen species (ROS and RNS), such as hydrogen peroxide (H2O2), superoxide radical (O2,), hydroxyl radical ('OH), nitric oxide ('NO), and nitrogen dioxide radical

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('NO₂), leading to the decomposition of luminol and the formation of 3-aminophthalate, which emits light. 10-13 Additionally, certain catalysts, including peroxidase, 14,15 hemoglobin, 16 hemin (Fe(III)-protoporphyrin), 17,18 and enhance these oxidation reactions, significantly increasing luminescence intensity. Moreover, the CL quantum yield of the different luminol derivatives depend on the solvent polarity, nature, position and number of substituents. 19,20

To enhance the applicability of luminol, researchers have focused on improving its sensitivity, stability, and solubility through the synthesis of more potent chemiluminescent derivatives. This concise review is structured into three main sections (Fig. 1). The first section provides an overview of luminol synthesis, its structural features and functional groups, the underlying CL reaction mechanisms, and key factors influencing luminescence efficiency. The second section explores various modification strategies for luminol derivatives, including structural alterations to enhance sensitivity, stability, and solubility, functionalization with electron-donating (EDG) or withdrawing groups (EWG), and the development of hybrid luminol derivatives with superior properties. The final section examines challenges and future perspectives, focusing on stability and storage limitations, the need for more biocompatible and selective derivatives, emerging synthesis trends, and potential interdisciplinary applications.

2. Chemistry of luminol and mechanism of luminescence generation

2.1. Luminol synthesis approaches

Luminol synthesis begins with 3-nitrophthalic acid, which can be obtained via the oxidation of α -nitronaphthalene or



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with particular emphasis on nitric oxide and its role in cancer and inflammation. Dr. Alsharabasy has published extensively on chemiluminescent sensors, gasotransmitter-responsive biomaterials, and redox-based therapeutics. His work integrates synthetic chemistry and materials science to advance next-generation diagnostic and therapeutic platforms for biomedical use.

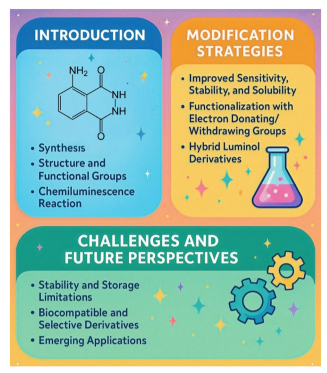


Fig. 1 Overview of the review structure, including luminol synthesis, chemiluminescence reaction mechanism and light generation, strategies for luminol derivative design, and future challenges and applications.

the nitration of phthalic acid or phthalic anhydride (Scheme 1). The synthesis starts with naphthalene nitration, which can be performed using: (1) nitric acid at room temperature for 5-6 days, followed by purification via methanol recrystallization.21 (2) Sulphonitric acid at 50-60 °C for 7 hours, followed by purification with xylene.²² (3) Sulphonitric acid at 45-60 °C for 90 minutes, followed by purification with diluted ethanol.²³ (4) Sulphonitric acid at 95 °C, nitrating molten naphthalene for 30 minutes, followed by purification with methanol (step 1, Scheme 1).²¹

The resulting α -nitronaphthalene is then oxidized by mixing with glacial acetic acid and chromic anhydride, heating at 70-80 °C for 90 minutes, followed by cooling, filtration of insoluble impurities, and washing.21 The solution is subsequently treated with barium carbonate and sodium carbonate, boiled, and then acidified with HCl at 50-60 °C to yield 3-nitrophthalic acid (step 2, Scheme 1). The nitration of phthalic acid or phthalic anhydride using Culhane's method produces a mixture of 3-4-nitrophthalic acids, which are difficult to separate.²⁴

To obtain 3-nitrophthalhydrazide, 3-nitrophthalic acid is mixed with hydrazine sulphate and allowed to evaporate to dryness, followed by heating with decalin at 160 °C for 3 hours.²⁵ The cooled product is then precipitated with benzene, filtered, and washed with ethyl ether (step 3, Scheme 1). Alternatively, 3-nitrophthalic acid can be mixed with a hydrazine solution in triethylene glycol, refluxed at $^{\circ}\mathrm{C}$ for 2 minutes, and cooled yield 3-nitrophthalhydrazide.²⁶ In the final stage,

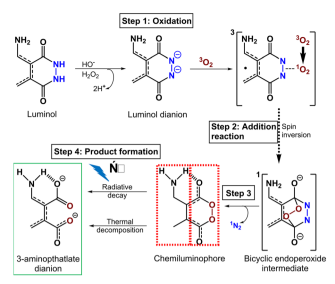
Scheme 1 Luminol synthesis pathways, starting with naphthalene nitration (step 1), followed by oxidation to 3-nitro-phthalic acid (step 2), which, upon hydrazinolysis (step 3), and subsequential treatment with NaOH, sodium dithionite, and acidic precipitation yield luminol (step 4).

3-nitrophthalhydrazide is mixed with sodium sulphate and NaOH or NH4OH, followed by the addition of sodium dithionite and heating at 70 °C for 30 minutes.²⁷ The solution is then acidified with glacial acetic acid, and the precipitate is washed with water, dissolved in hot HCl, briefly decolorized with charcoal, filtered, washed, and dried (step 4, Scheme 1).

2.2. Mechanism of luminescence generation

Light emission in luminol solutions occurs under alkaline conditions in the presence of oxygen, following specific mechanisms in both protic and aprotic media.²⁸ Scheme 2 summarizes these CL reactions. In aprotic solutions, the luminol dianion forms and facilitates the spin inversion of molecular oxygen from its triplet state (${}^{3}O_{2}$) to singlet oxygen (1O2).29 This leads to a concerted addition reaction, forming a bicyclic endoperoxide intermediate. The nitrogen atoms are then replaced by oxygen atoms through a peroxide bond, resulting in the formation of a chemiluminophore and the release of nitrogen gas. Electron transfer from the aniline ring to the O-O bond enables either high-energy or lowenergy chemiexcitation. This process involves cleavage of the -O-O- bond, followed by either radiative decay (light emission) or radiationless decay.

The emission wavelength in aprotic solvents (e.g., DMSO) typically ranges from 485 to 510 nm. In protic solvents (e.g., water), the emission undergoes a blue shift to 420-430 nm due to intramolecular hydrogen transfer from the amino group to a nearby oxygen atom. 30-32 The duration of light emission varies from a few seconds to several hours, depending on factors such as the medium composition,³³ the reacting species,34 and the presence of additives or coinclude peroxidases,35 These various metalloporphyrins,36 ROS and RNS.13



Scheme 2 Chemiluminescence of luminol in aprotic solvents, starting with the oxidation to luminol dianion (step 1), followed by addition reaction and spin inversion of molecular oxygen (step 2). Upon nitrogen elimination and formation of chemiluminophore (step 3), 3-aminophthalate is formed via chemiexcitation, followed by radiative or non-radiative decay (step 4).

3. Advances in the synthesis of luminol derivatives

Despite its wide range of applications, luminol has several limitations, including poor solubility in aqueous solutions, short luminescence duration, and weak intensity under certain conditions. To address these drawbacks, various chemical modification strategies have been developed, primarily focusing on introducing new functional groups to the C-NH₂ position, 37,38 and substitutions on the phenyl ring to modulate electron-donating effects. 20,39 These substituent effects generally alter the electron density, highest occupied molecular orbital (HOMO), and lowest unoccupied molecular orbital (LUMO) energy levels, pK_{α} , and redox potentials. For instance, the conjugation of luminol to EDG is expected to stabilize the luminol dianion resonance, reducing the energy penalty for deprotonation, 19 which consequently make the CL reaction more feasible at physiological pH (7-8) instead of requiring strong alkali (pH ≥ 10). While these modifications generally aim to improve CL efficiency, some substitutions at the C-NH₂ position have been found to reduce luminescence intensity by disrupting proton transfer between the amino and carbonyl groups, which is essential for light emission.²⁹ Moreover, the luminol conjugation to EWG (e.g., chloro and nitro groups) reduced the excitedstate stability, with a general decrease in luminescence intensity.40,41 Additionally, the chloride derivatives of mefenamic acid, ibuprofen, diclofenac sodium, and ampicillin were reacted with luminol in DMSO and triethylamine for the development of a new class of luminol derivatives with antibacterial activity. 42 However,

the luminescence activity of these derivatives was not This investigated further. section summarizes key advancements in the development of luminol derivatives with enhanced luminescence properties, highlighting the characteristics of promising candidate compounds.

3.1. Substitutions on phthalimides

Mikroulis et al. synthesized a series of 6,8-disubstituted derivatives using coupling luminol Suzuki hydrazinolysis.43 The synthesis began with the refluxing of

Scheme 3 (A and B) Synthesis route of the 3-amino-phthalimide, followed by the synthetic pathways of 6-methyl-8(4-hydroxymethylphenyl) luminol and 6,8-dimethyl luminol, respectively. (C) Synthesis route of phosphonium phthalhydrazide beginning with the reaction between 3-aminophthalimide and bromoalkyl carboxylic acids for phthalimide acylation, followed by synthesis of phosphonium phthalimide, and final hydrazinolysis. Abbreviations: 4-HBPA: 4-(hydroxymethyl)phenylboronic acid; TMB: trimethylboroxine.

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3-nitro-phthalic acid and acetic anhydride (Ac₂O), followed by solvent evaporation in the presence of toluene and precipitation of the anhydride upon diethyl ether addition.³⁸ The resulting anhydride was refluxed then sec-butylamine and acetic acid, dried, dissolved in dichloromethane (DCM), and washed with an aqueous NaHCO3 solution. After drying with Na2SO4, this process yielded 3-nitro-phthalimide. The nitro group was subsequently reduced to an amino group by degassing the 3-nitro-phthalimide in methanol, adding Pd/C, and exposing the mixture to H₂ gas, forming 3-amino-phthalimide. Bromination was carried out using either excess N-bromosuccinimide in DCM or bromine with acetic acid and sodium acetate, vielding 6-monobrominated 4,6-dibrominated derivatives, respectively (Scheme 3A). Suzuki methylation with trimethylboroxine produced the corresponding 6-monosubstituted and 4,6-disubstituted 3-aminophthalimide derivatives.

enhance luminescence intensity the To 6-monosubstituted derivative, an additional bromination step was performed, followed by reaction with 4-(hydroxymethyl) phenylboronic acid to introduce a hydroxyl-functionalized group (Scheme 3B). Hydrazinolysis of these products yielded luminol derivatives with enhanced CL efficiency under biologically relevant conditions. Notably, the CL quantum of 6,8-dimethyl luminol and 6-methyl-8-(4hydroxymethylphenyl) luminol were 0.17 respectively, at pH 12, compared to just 0.012 for luminol (Table 1). Importantly, the enhanced luminescence observed with 8-methyl substitution compared to 8-phenyl is attributed to its ability to donate negative charge to the cyclic peroxide, facilitating O-O bond cleavage. Substitution at position 6 has a smaller impact on this effect. Similar enhancements in luminescence intensity were observed with luminol derivatives containing a 6-monomethyl substitution, which exhibited a twofold increase in CL intensity compared to luminol.44 However, no significant differences were noted for propyl- or tert-butyl-substituted derivatives, as the steric interactions of these alkyl groups with the carbonyl group hindered the transition to the excited state.

Interestingly, phosphonium cations were introduced into luminol derivatives to enable mitochondrial targeting for detecting intracellular ROS.38 Phosphonium-functionalized amino-acylated luminol derivatives were synthesized by stirring bromoalkyl carboxylic acids with oxalyl chloride (COCl)₂ under an argon atmosphere, followed by evaporation of volatiles, redissolution in DCM, and reaction with 3-amino-phthalimide in the presence of pyridine (Scheme 3C). The resulting product was purified via aqueous two-phase extraction using water and DCM, followed by drying with Na2SO4. The acylated product was then reacted with phosphine in dry acetonitrile and refluxed under argon for three days, yielding the phosphonium phthalimide, which was subsequently refluxed with hydrazine in ethanol for hydrazinolysis. Although these final mitochondriotropic chemiluminescent probes successfully targeted mitochondria, they exhibited lower luminescence efficiency than luminol. Their poor CL performance and reduced intensity (Table 1) were attributed to fluorescence inhibition during the oxidation reaction. A similar reduction in luminescence intensity was observed after modifying isoluminol using the same approach.

Another approach for synthesizing luminol derivatives with substitutions at positions 6 and 8 was reported before, 45,46 as shown in Fig. 2A. These studies aimed to modulate the steric interactions between the introduced alkyl groups and the

Table 1 Comparative summary of luminol derivatives: structural modifications, quantum yields, and emission wavelengths

Luminol derivative	Structural modification	Quantum yield	Emission wavelength (nm)	Ref.
Luminol	Parent compound	0.012	425	44
6,8-Dimethyl luminol	Methyl substitutions at positions 6 and 8 on the benzene ring	0.17	425	44
6-Methyl-8-(4-hydroxymethylphenyl) luminol	Methyl group at position 6 and 4-hydroxymethylphenyl at position 8	0.39	425	44
Phosphonium phthalhydrazide	Phosphonium group substitution on the phthalhydrazide structure	< 0.001	425	38
6-Methyl luminol	Single methyl substitution at position 6	0.237	425	47
6-Isopropyl luminol	Isopropyl group at position 6	0.012	425	47
Naphthalene-based analogues (GL-1 to GL-4)	Benzene ring replaced by naphthalene to extend conjugation	0.3 (GL-1)	520	50
Luminol-fluorescein based activated cassettes	Luminol chemically linked to fluorescein <i>via</i> a linker	0.007	524	52
Luminol-Nile red based activated cassettes	Luminol chemically linked to Nile red via a linker	0.012	634	52
Isoluminol	Amino group at the <i>meta</i> -position	≈0.0012	416	41
O-Etherified luminol (Lum-glucose)	Enol group etherified with glucose <i>via</i> O-linkage	Quenched (restored after β-glu cleavage)	Turn-on signal after enzyme cleavage	53
m-Carboxyl luminol	Carboxyl group introduced at the <i>meta</i> -position	0.06		39
Acylated luminol hydrazides	Succinic/glutaric acid linkers on the amino group	0.004	Minor shifts	75

maleimide

Fig. 2 (A) Schematic representation of the synthesis of alkyl luminol derivatives. (B) Synthetic routes of luminol derivatives with naphthalene, and the corresponding CL emission spectra in $H_2O/DMSO$ solution with potassium persulfate in the inset. Reprinted with permission.⁴⁹

2 and 3

carbonyl group of luminol. The synthesis began with the coupling of *O*-benzyl carbamate, *N*-methyl maleimide, and a series of unsaturated aldehydes, yielding a group of protected aminocyclohexenes. Subsequent deprotection using palladium-on-charcoal, followed by hydrazinolysis, produced a series of 6,8-dialkyl luminol derivatives. Among the synthesized derivatives, 6-methyl luminol exhibited a 20-fold increase in luminescence efficiency compared to luminol,

twofold accompanied by enhancement in chemiluminescence quantum yield and a slight blue shift (Table 1). In contrast, 6-isopropyl luminol displayed luminescence properties like those of luminol.47 These studies highlighted the role of steric effects introduced by alkyl substituents, which facilitate the transition to the excited state and enhance CL yields. This steric-based mechanism complements the previously discussed

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electronic modifications and suggests the potential of these luminol derivatives for bioluminescence imaging.

Notably, some derivatives demonstrated solubility in both aqueous and aprotic solutions, broadening their applicability. However, the synthesis of certain derivatives still requires long reaction times and palladium catalysis, which is both costly and highly sensitive to reaction conditions. Further optimization is needed to improve the efficiency and scalability of these synthetic routes.

Another strategy for enhancing luminol's luminescence involved extending its aromatic system by replacing the benzene ring with a naphthalene derivative functionalized with an -NH2 group or further modifying this exocyclic group. The synthesis began with the hydrazinolysis of 2,3-naphthalenedicarboxylic acid in acetic acid, followed by nitration using nitronium tetrafluoroborate (BF4NO2) in acetonitrile and subsequent mild reduction of the -NO2 group at position 5 (Fig. 2B). 48,49 For derivatives with a protected -NH₂ group, the reaction pathway started with esterification of 2,3-naphthalenedicarboxylic acid, followed by nitration and reduction. This was followed by monoand di-benzylation of the amine group in the presence of KHCO3 or NaOH in DMF, with final hydrolysis and hydrazinolysis steps. Additional derivatives featuring amine substitution at position 6 were also synthesized, though they are not shown here. These naphthalene-based luminol derivatives exhibited green luminescence within the 490-590 nm range, with longer decay times and stronger CL signals than luminol (the inset in Fig. 2). As a result, they were proposed for the detection of lowconcentration analytes, offering improved sensitivity compared to luminol.

Another strategy involved the synthesis of green-emitting luminol analogues through a three-step reaction sequence starting from 1-naphthylamine, including diazotization, nucleophilic substitution, and cyclization (Fig. 3A). This synthetic modification replaced luminol's benzene core with a naphthalene moiety, effectively extending the π -conjugation

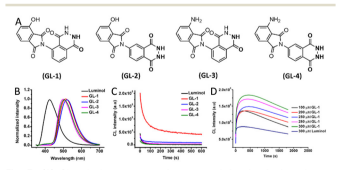


Fig. 3 (A) Chemical structures of green-emitting luminol analogues (GL-1 to GL-4). (B) Luminescence emission spectra comparison between luminol and its analogues. (C) CL kinetic profiles of luminol and GL analogues. (D) CL intensity over time for varying concentrations of GL-1 and luminol in the presence of HRP, hydrogen peroxide H₂O₂, 3-(10'-phenothiazinyl)-propane-1-4-morpholinopyridine, and sulfonate.53

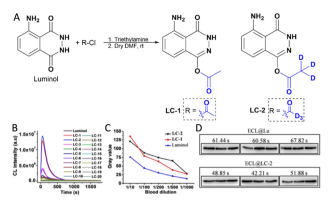


Fig. 4 (A) Schematic representation of the synthesis of 5-amino-4oxo-3,4-dihydrophthalazin-1-yl acetate (LC-1) and its deuterated counterpart (LC-2). (B) CL response curves, showing intensity versus time for luminol and its derivatives in the presence of H2O2 and hemin as a catalyst. (C) Gray value analysis of bloodstain images captured using a smartphone, following detection using a CL solution containing luminol, LC-1, or LC-2. (D) Western blot (WB) analysis of GAPDH protein expression, detected in mixtures containing luminol or LC-2 with H₂O₂. Reprinted with permission.⁵¹

and red-shifting the emission wavelength to approximately 520 nm (Fig. 3B). In addition to the spectral shift, the structural changes enhanced the compound's stability under ambient conditions. Among the synthesized derivatives, the analogue GL-1 exhibited the highest CL yield compared to both luminol and other analogues (Fig. 3C), intensity correlating linearly with luminescence concentration (Fig. 3D, Table 1).50

3.2. Luminol modification via O-esterification and Oetherification

To enhance the CL intensity of luminol, an O-esterification reaction was performed by mixing it with triethylamine in DMF at room temperature or 50 °C (Fig. 4A). 51 Among the synthesized derivatives, 5-amino-4-oxo-3,4-dihydrophthalazin-1-yl acetate (LC-1) and its deuterated counterpart LC-2 exhibited the highest luminescence efficiency, with significantly improved intensity and prolonged emission time compared to luminol and other derivatives within the pH range of 9 to 11. However, no details on the quantum yields were reported. Representative results from mixtures containing H₂O₂ and hemin at pH 9 are shown in Fig. 4B. Notably, reactions containing LC-2, which features a deuterated acetyl group, exhibited the highest reaction rate in the presence of blood (Fig. 4C). This enhancement is attributed to the esterification of the hydroxyl group, which stabilizes luminol in its enol form, thereby increasing luminescence intensity. Additionally, both luminol and LC-2 demonstrated comparable luminescence efficiency in western blot (WB) imaging for GAPDH detection, using goat antimouse IgG (H + L)-HRP as the secondary antibody (Fig. 4D). However, further esterification with larger substituents may negatively impact solubility and, due to steric hindrance, could reduce luminescence efficiency (Fig. 2B). These steric

effects likely explain the lower performance observed in other synthesized derivatives.

Development of luminescence-based probes biotechnological applications emitting at wavelengths above 450 nm has shown some success. For example, Han et al. studied chemiluminescent energy-transfer cassettes by coupling luminol derivatives with fluorescein and Nile red via Sonogashira reactions. 52 The luminescence yield of luminol-fluorescein and luminol-Nile red was 61% and 100%, respectively, relative to luminol, producing yellow/ green and red CL (Table 1). However, despite the efficient energy transfer, the synthesis involved multiple steps, lacked high convergence, and resulted in poor solubility in aqueous media, limiting biological applications. Additionally, these derivatives lack the amino group of luminol, which affect their CL efficiency and reactivity. Furthermore, switchable luminol-based electro-chemiluminescence (ECL) biosensors were developed by O-etherifying luminol at the enol group and conjugating it to glucose *via* an ether bond (Table 1).⁵³ The resulting conjugate exhibited suppressed ECL emission, which was restored upon enzymatic cleavage by β-glucosidase, enabling sensitive detection of the enzyme with a linear range of 0.4-400 U L⁻¹ and a detection limit of $0.1~{\rm U}~{\rm L}^{-1}$.

3.3. Approaches for enhancing the solubility of luminol

Despite luminol's wide applicability in various analytical approaches, its poor solubility in aqueous solutions remains a challenge for specific assays. To address this, several

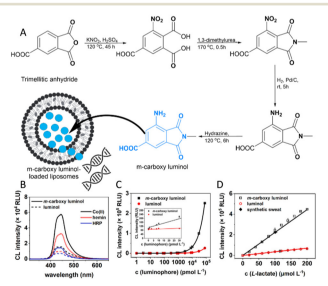


Fig. 5 (A) Schematic representation of the m-carboxy luminol synthesis. (B) CL response curves for luminol and m-carboxy luminol with H_2O_2 in the presence of cobalt chloride (Co^{2+}), hemin, and horseradish peroxidase (HRP) as catalysts. (C) CL response curves for varying concentrations of luminophores (luminol and m-carboxy luminol) mixed with H₂O₂ and hemin. (D) CL response curves corresponding to different lactate concentrations. Data are presented as mean \pm SD (n = 3). Reprinted with permission.³⁹

studies have explored the synthesis of more hydrophilic luminol derivatives. For example, introducing a carboxyl (-COOH) group into luminol's benzene ring was achieved through the nitration of trimetallic anhydride at 120 °C, followed by treatment with dimethylurea, reduction of the exocyclic nitro group to an amine, and a final hydrazinolysis step (Fig. 5A). 54,55

The resulting *m*-carboxyl luminol exhibited improved water solubility and, when loaded into liposomes, produced a fourfold increase in luminescence intensity with a 150-fold lower DNA detection limit compared to luminol.⁵⁵ Moreover, in the presence of cobalt chloride, hemin, and HRP, m-carboxyl luminol exhibited a fivefold increase in quantum yield compared to luminol, along with ~18-fold higher sensitivity for H₂O₂ detection (Fig. 5B, Table 1).³⁹ Additionally, the lower detection limit for hemin was reduced fivefold with m-carboxyl luminol compared to luminol (Fig. 5C). Similarly, the luminescence signal in response to lactate was 2.6 times higher with m-carboxyl luminol, with generally lower detection limits than those obtained using colorimetric measurements (Fig. 5D). While this derivative shows potential for integration into microfluidic systems for point-of-care diagnostics, its low yield (15%) requires further optimization.⁵⁵

3.4. Macromolecular strategies for luminol functionalization and signal amplification

Conjugating luminol to the backbones of various macromolecules has been employed to unlock a range of chemical, physical, and application-specific advantages. This strategy aims to achieve the following objectives: (1) enhanced stability of the chemiluminophore through covalent attachment, which helps prevent leaching and degradation, thereby increasing the overall longevity of the system.⁵⁶ (2) Customizable chemical design that enables tuning of the chemical structure of redox-active materials by controlling the local environment around luminol to achieve specific functional properties. The main polymerization methods for luminol include oxidative chemical polycondensation, microwave-assisted oxidative polymerization, free-radical polymerization, and other miscellaneous approaches, which have been described in detail previously.⁵⁷ (3) Reusability, where polymer-bound luminol allows for reagentless and repeatable detection, as demonstrated various electropolymerized in sensors derivatives.58-64 (4)incorporating luminol Signal amplification, as multiple luminol units can be attached to a polymer chain, significantly boosting chemiluminescent output.⁵⁷ (5) Multifunctionality, enabling the incorporation of luminol alongside other functional groups (e.g., targeting moieties) on the same polymer for advanced biosensing applications.^{65,66} (6) Improved physical properties of the final luminol-based materials, where the inherent mechanical strength, processability, and thermal stability of polymers allow luminol to be applied in durable

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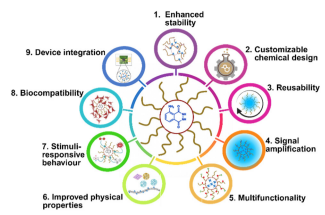


Fig. 6 Main objectives for conjugating luminol to the backbones of macromolecules.

forms such as films, fibres, and nanoparticles.⁵⁷ (7) Stimuli-responsive behaviour, where luminol-polymer conjugates act as self-reporting materials that emit light in response to specific environmental triggers such as oxidants, pH changes, or target analytes.^{58,67} (8) Biocompatibility, achieved by using suitable polymer carriers that enhance luminol's solubility and compatibility with biological fluids, enabling its use in biomedical applications.^{65,66} (9) Device integration, wherein luminol-polymer conjugates facilitate the incorporation of luminol into various electronic and optical systems, showing promise for embedded light-emitting functionalities.^{68,69} These advantages are summarized in Fig. 6. The recent advances in the development of luminol based polymers have been reviewed before.⁵⁷

Various strategies for loading of branched polymers with different luminophores towards achieving certain fluorescence properties and certain applications have been developed.⁷⁰ Dendrimers, such as poly(amidoamine) (PAMAM) and poly(dithiodipropionic acid) (PDTPA), have been strategically integrated into luminol-based ECL biosensor platforms to enhance performance through multiple synergistic Their highly branched three-dimensional mechanisms. architecture provides abundant terminal groups, enabling dense luminol loading for multi-labelling and significant signal amplification. For instance, a luminol-PDTPA-Fe₃O₄ nanoparticle-based immunosensor was developed for the detection of carbohydrate antigen 125 (CA125) using a sandwich structure consisting of a primary antibody and a luminol-labeled secondary antibody via a PDTPA dendrimer. This complex was anchored to Fe₃O₄ magnetic nanoparticles and immobilized on a carbon paste electrode through magnetic force, yielding a detection range of 0.2-100 mU mL⁻¹ and a detection limit of 0.032 mU mL⁻¹.71 A similar strategy was applied for the detection of microalbuminuria-a key marker for early-stage renal complications in diabetes and hypertensive pregnancy disorders—using a microalbuminuriaspecific antibody in a luminol-dendrimer-Fe₃O₄ system. This sensor achieved a detection range of 2-322 pg mL⁻¹ and a detection limit of 0.084 pg mL⁻¹.⁷²

Scheme 4 (A) Synthesis routes of luminol derivatives containing terminal hydrazide groups, using succinic acid and glutaric acid as linker moieties. (B) HRP labelling with the synthesized luminol-amido- C_2 -hydrazide, following enzyme oxidation with sodium periodate.

In addition to enabling high luminol loading, dendrimers act as efficient scaffolds for bioconjugation, facilitating the co-attachment of luminol and recognition biomolecules while enhancing stability by minimizing luminol leaching or degradation. For example, PAMAM-functionalized ZnO nanorods were employed for the simultaneous immobilization of luminol and a detection antibody specific to cancer antigen 15-3 (CA15-3). The ZnO nanorods catalyzed the decomposition of hydrogen peroxide, generating reactive oxygen species (ROS) that triggered and amplified the ECL signal. This system enabled CA15-3 quantification with a dynamic range of 0.1-120 U mL⁻¹ and a detection limit of $0.033~U~mL^{-1}$.73

For the development of advanced CL immunoassays, labelling functional enzymes (*e.g.*, peroxidase, phosphatase) with a CL-labelling molecule would be advantageous. However, compared to fluorescence-labelling reagents, the introduction of CL molecules remains challenging, as many are chemically sensitive, and modifications can significantly affect luminescence intensity. ^{30,74} For instance, acylation of the exocyclic amino group in luminol was achieved by overnight reaction with either succinic or glutaric anhydride in alkaline water, followed by hydrazinolysis in the presence of 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC) (Scheme 4A). ⁷⁵

4. Challenges and future perspectives

Stability, storage, and biocompatibility limitations

Despite the promise of luminol-based systems, key translational barriers remain. One of the most critical limitations is the instability of luminol derivatives under ambient conditions. Compounds such as hydrazides are prone to autoxidation and hydrolysis, while others suffer from photobleaching, collectively diminishing CL quantum yields within weeks.⁷⁶ Current storage protocols, such as freezing at -20 °C under an inert atmosphere and light

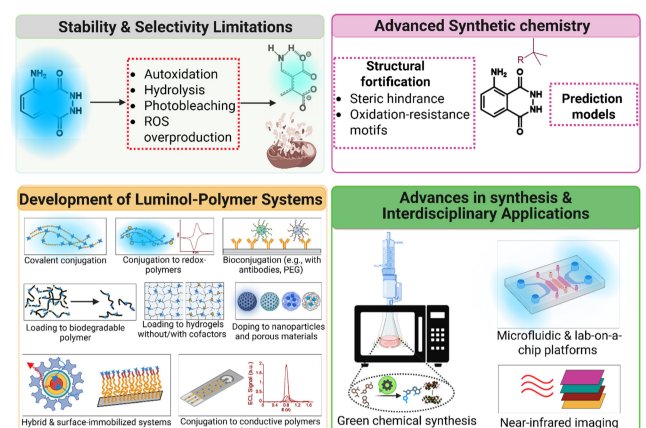


Fig. 7 Challenges and emerging strategies in luminol-polymer chemiluminescent systems.

exclusion, are effective but impractical for field deployment or point-of-care applications.

Equally important are biocompatibility and selectivity challenges, which limit in vivo utility. Unmodified luminol derivatives may cause dose-dependent cytotoxicity due to excessive ROS production and mitochondrial disruption.^{77,78} Additionally, nonspecific interactions with serum proteins or metal ions can hinder signal specificity and reduce assay accuracy.⁷⁹ These limitations are summarized in Fig. 7.

Advanced synthetic chemistry approaches

To overcome these challenges, future innovations should prioritize chemical strategies that enhance molecular robustness and predictive stability. These include: (1) structural fortification through sterically hindered N-alkyl groups (e.g., tert-butyl) and oxidation-resistant motifs (e.g., thiadiazole analogues) to prevent degradation.^{80,81} Machine learning-guided design to develop predictive models capable of identifying luminol derivatives with long-term shelf stability (Fig. 7).

Development of advanced luminol-polymer systems

Polymer-based platforms provide a promising avenue to address the inherent weaknesses of free luminol, including short emission duration, pH dependency, and lack of targeting. These strategies not only amplify the signal but also introduce spatial control, enhanced stability, and multifunctionality (Fig. 7).

& computational chemistry

The first strategy depends on covalent functionalization of polymers, where grafting luminol onto polymers (e.g., those with carboxylate groups) helps mitigate pH sensitivity, improve aqueous solubility, and support physiological performance. Branched architectures, such as dendritic polymers, offer steric shielding,82 which, upon conjugation to luminol, can reduce quenching, increase its local density, and enhance CL intensity in biosensing platforms. Redoxactive polymers (e.g., poly(vinylferrocene)) offer alternative luminol activation pathways for self-powered sensing. Bioconjugation with PEG, peptides, or antibodies/aptamers enhances biocompatibility, reduces immune clearance, and enables target-specific recognition.83 Additionally, using biodegradable polymers (e.g., PLGA) allows for safe degradation and reduces cytotoxicity.84

The second strategy spans a number of encapsulation approaches, where hydrogel-based luminol entrapment provides structural protection and spatial control. Coencapsulation with catalytic cofactors such as hemin, horseradish peroxidase (HRP), or heme proteins enables reagent-free, enzyme-driven luminescence suitable for pointof-care detection.85-88 Our recent work with hemin-loaded HA hydrogels shows promise for NO detection, and ongoing Review **MSDE**

efforts aim to incorporate luminol directly into the matrix for simplified fabrication and enhanced reagent stability.89 Additionally, stimuli-responsive systems activated by diseaserelated enzymes (e.g., MMPs or MPO) enable site-specific CL for real-time biomarker monitoring. Moreover, encapsulation within protective nanomaterials can improve luminol's stability and quantum yield. Silica nanoparticles doped with luminol have demonstrated improved performance, 90-92 though further development is needed. Similarly, luminolloaded metal-organic frameworks (MOFs) have been employed as chemiluminescent scaffolds in biosensing. 93-95

The third strategy is the development of hybrid and surface-immobilized systems. Multifunctional materials such as surface-immobilized polymer brushes and luminolquantum dot micelles combine CL with resonance energy transfer (CRET) for tuneable biosensing. 96,97 Conductive polymer conjugates also allow for low-voltage ECL by serving as both the catalyst and the electrode. These platforms offer highly modular, reagent-free solutions to conventional luminol limitations.

Advances in synthesis and interdisciplinary applications

Scalability and environmental sustainability are critical in luminol derivative synthesis. Recent trends emphasize green chemistry, such as microwave-assisted polymerization, which reduces reaction time and energy consumption. 98,99 Aqueousphase cross-coupling reactions (e.g., Suzuki-Miyaura) offer solvent-free alternatives for luminol functionalization. ¹⁰⁰ Ecofriendly catalysts and catalyst recycling further support sustainable practices.

In parallel, computational chemistry, including density functional theory (DFT) and AI-based molecular design, can accelerate the discovery of new luminol derivatives by predicting quantum yields optimizing chemiluminescence profiles. 101,102

derivatives Luminol are also extending interdisciplinary domains such as bioimaging, point-of-care diagnostics, and environmental monitoring. For instance, near-infrared (NIR) luminol analogues are being investigated for deep-tissue imaging, and multi-layer films generating long-lasting NIR emission via peroxyoxalate reactions may be adapted for these systems. 103 Integration with microfluidic and lab-on-a-chip platforms further enhances luminol's diagnostic utility. 104-106

Together, these emerging strategies position luminolpolymer systems as a next-generation platform for biosensing and diagnostic technologies. Future research should focus on scalable synthesis, AI-guided molecular engineering, and integration with biocompatible, degradable materials to enable real-world applications in clinical, environmental, and industrial settings.

Conclusions

Over the years, significant progress has been made in the synthesis of luminol derivatives, resulting in improved CL efficiency, enhanced solubility, and greater Structural modifications such as alkylation, esterification, and heterocyclic substitution have allowed precise tuning of luminol's photophysical properties, supporting its successful integration into diverse analytical platforms. Additionally, the incorporation of luminol into advanced hybrid materials, such as nanocomposites, polymer matrices, and metalorganic frameworks, has significantly expanded its functional range for biomedical diagnostics, forensic analysis, and environmental sensing.

Despite these advancements, critical challenges remain in translating luminol-based systems into real-world applications. Instability under ambient conditions. biocompatibility concerns, and lack of target specificity continue to limit in vivo and point-of-care deployment. To overcome these barriers, recent research has prioritized robust synthetic strategies, including sterically protected molecular designs and machine learning-guided enhance stability optimization, to and predictive performance. Polymer conjugation and nanoscale encapsulation approaches are also being leveraged to amplify signal output, extend luminescence duration, and introduce stimuli-responsive behaviour for dynamic sensing environments.

Looking forward, the development of scalable, green microwave-assisted synthesis methods, such as polymerization and aqueous-phase cross-coupling reactions, will he critical for sustainable manufacturing. Simultaneously, computational chemistry and artificial intelligence offer powerful tools for predicting and tailoring luminol's chemiluminescence behaviour. The continued convergence of synthetic innovation, materials science, and bioengineering will drive the evolution of luminol-based systems into next-generation diagnostic and sensing platforms. With interdisciplinary collaboration and a focus clinical and environmental translation, derivatives are poised to remain indispensable tools in analytical science and beyond.

Data availability

No new datasets were generated in this study. All data discussed are available in the published literature and cited accordingly.

Conflicts of interest

There are no conflicts to declare.

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