



Carcinogenesis

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Abstract

The relentless progression of carcinogenesis, characterized by uncontrolled cellular proliferation and the acquisition of metastatic potential, remains a predominant cause of global mortality. Conventional therapeutic modalities—chemotherapy, radiation, and surgery—often face insurmountable barriers including multidrug resistance, lack of specificity, and debilitating systemic toxicity. This research article presents an exhaustive pharmacognostic investigation of *Ganoderma lucidum* (Reishi), a Basidiomycete fungus historically revered for its life-extending properties, and positions it as a pivotal bio-resource in the green synthesis of a novel trimetallic nanotherapeutic system.

Our study rigorously authenticates *G. lucidum* through macroscopic, microscopic, and physicochemical analyses, confirming the presence of bioactive triterpenoids and polysaccharides capable of reducing metal salts. Leveraging this reducing capacity, we synthesized a hybrid core-shell nanoparticle architecture comprising superparamagnetic Iron Oxide (Fe₃O₄), plasmonic Gold (Au), and bioactive Selenium (Se). This investigation details the synthesis protocols, physicochemical characterization, and the theoretical underpinnings of utilizing this trimetallic system for simultaneous magnetic targeting, photothermal ablation, and selenium-mediated apoptosis. The results suggest that the integration of ancient pharmacognosy with avant-garde nanotechnology offers a paradigm shift in oncology, moving towards "green" theranostics that are both biocompatible and lethally effective against neoplastic cells.

Keywords

- Carcinogenesis
- Carcinoma
- genes
- Expressions
- Phenotype
- Genotype
- Drug Response
- *Ganoderma lucidum*
- Gold
- Selenium
- Iron oxide

INTRODUCTION

The Global Burden of Carcinogenesis

Carcinogenesis is not a singular event but a complex, multi-stage evolutionary process wherein normal somatic cells undergo neoplastic transformation. This transformation is driven by the accumulation of genetic and epigenetic alterations that disrupt the delicate balance between cell division and programmed cell death (apoptosis). The burden of this disease is staggering; it represents a primary barrier to increasing life expectancy in every nation of the world in the 21st century. Despite decades of intensive research, the heterogeneous nature of tumors and their ability to evolve resistance to therapies continue to confound standard treatment protocols.

The pathogenesis of cancer involves three distinct phases: initiation (irreversible genetic damage), promotion (clonal expansion of initiated cells), and progression (acquisition of malignant features such as invasiveness and metastasis). Conventional chemotherapeutics typically target rapidly dividing cells, a strategy that lacks precision and invariably damages rapidly dividing healthy tissues—bone marrow, gastrointestinal epithelium, and hair follicles—leading to the severe side effects that limit dosage and efficacy. Furthermore, the tumor microenvironment (TME) is often hypoxic and acidic, conditions that can neutralize drug efficacy and promote the selection of resistant cell clones.

The Limitations of Current Therapeutics

The current standard of care faces a "therapeutic ceiling." Radiation therapy is limited by the tolerance of surrounding normal tissues. Surgical resection is often impossible in metastatic or hematological malignancies. Chemotherapy is plagued by the phenomenon of Multidrug Resistance (MDR), mediated by efflux pumps like P-glycoprotein that actively expel drugs from cancer cells. Consequently, there is an urgent clinical imperative to develop "smart" therapeutic agents that can:

1. Selectively accumulate in tumor tissues (Active Targeting).
2. Deliver therapeutic payloads without systemic leakage (Controlled Release).
3. Overcome biological barriers and resistance mechanisms (Multimodal Action).
4. Provide real-time feedback on treatment efficacy (Theranostics).

The Renaissance of Pharmacognosy: *Ganoderma lucidum*

In the search for such agents, modern science is increasingly revisiting the pharmacopoeia of antiquity. *Ganoderma lucidum*, known as "Lingzhi" in China and "Reishi" in Japan, stands at the apex of this revival. Unlike pharmaceutical agents synthesized de novo, *G. lucidum* has co-evolved with biological systems, producing a diverse array of metabolites—triterpenoids, polysaccharides, and proteins—that modulate mammalian signaling pathways with high specificity.

Historically, *G. lucidum* was reserved for nobility and believed to bestow immortality. Pharmacognostically, it is defined not just as a folk remedy but as a reservoir of phytochemicals with proven cytotoxic, immunomodulatory, and hepatoprotective activities. The dual capacity of this mushroom to directly attack cancer cells (via triterpenoids) and simultaneously bolster the host immune system (via polysaccharides) makes it a unique candidate for integrative oncology. However, the bioavailability of crude extracts is often low, necessitating advanced delivery systems.

The Convergence: Green Nanotechnology

Nanotechnology offers the solution to bioavailability and targeting issues. Nanoparticles (NPs) ranging from 1 to 100 nm can exploit the Enhanced Permeability and Retention (EPR) effect, passively accumulating in the leaky vasculature of tumors. However, the chemical synthesis of NPs often involves toxic reducing agents (e.g., hydrazine, sodium borohydride) and organic solvents that render the final product biologically incompatible or require extensive purification.

"Green Synthesis" represents a sustainable alternative. By utilizing the phytochemical-rich extract of *G. lucidum* as both a reducing agent (to convert metal ions to metallic NPs) and a capping agent (to stabilize the NPs), we can produce biocompatible nanohybrids. This study explores the fabrication of a trimetallic system—Fe₃O₄/Au/Se—using *G. lucidum*. Each component plays a strategic role:

- **Iron Oxide (Fe₃O₄):** Provides superparamagnetism for MRI imaging and magnetic hyperthermia.
- **Gold (Au):** Offers surface plasmon resonance for photothermal therapy and CT contrast.
- **Selenium (Se):** Delivers direct chemotherapeutic toxicity via oxidative stress induction.

Pharmacognostic Study of *Ganoderma lucidum*

The foundation of any natural product research is the unambiguous identification and quality control of the source material. *Ganoderma lucidum* exhibits significant phenotypic plasticity, meaning its appearance can vary widely based on environmental conditions (substrate, temperature, humidity, CO₂ concentration). Therefore, a multi-tiered identification protocol involving taxonomic, macroscopic, and microscopic evaluation is essential to distinguish it from related species like *G. sinense* or *G. tsugae*.

Taxonomy and Botanical Description

Taxonomic Hierarchy:

- **Kingdom:** Fungi
- **Phylum:** Basidiomycota (Characterized by the production of basidiospores on a basidium).

- **Class:** Agaricomycetes (The mushroom-forming fungi).
- **Order:** Polyporales (Pored fungi, distinct from gilled Agaricales).
- **Family:** Ganodermataceae.
- **Genus:** Ganoderma.
- **Species:** Ganoderma lucidum (Curtis) P. Karst.

Ecological Niche: In the wild, *G. lucidum* is a white-rot fungus. It degrades lignin in wood, primarily growing as a saprophyte on decaying hardwood trees (oaks, maples, elms) or the base of stumps. It is widely distributed across the temperate and subtropical zones of Asia, Europe, and the Americas. The mycelium colonizes the wood, secreting enzymes like laccase and manganese peroxidase to break down complex lignocellulose structures.

Cultivation and Source Material: The source material for pharmaceutical use is rarely wild-harvested due to scarcity and inconsistency. Artificial cultivation is the standard, utilizing either:

1. **Wood-Log Cultivation:** Mimics natural conditions, yielding fruiting bodies with high triterpenoid content.
2. **Sawdust Bags:** Faster yield but potentially lower density of bioactive compounds.
3. **Submerged Fermentation:** Liquid culture used primarily to produce mycelial biomass and extracellular polysaccharides.

For this research, the pharmacognostic relevance of the maturity stage is critical. The "budding" stage is often richer in low-molecular-weight triterpenoids, whereas the mature fruiting body accumulates higher concentrations of high-molecular-weight polysaccharides and spores. The material used in this study represents the mature fruiting body to maximize the diversity of reducing agents available for nanoparticle synthesis.

Macroscopic Identification

Macroscopic evaluation—organoleptic analysis—provides the first tier of authentication. The unique visual characteristics of *G. lucidum* are so distinct that they serve as immediate diagnostics against gross adulteration.

Organoleptic Profile:

- **Appearance (Pileus):** The cap is kidney-shaped (reniform) to fan-shaped (flabelliform). The most striking feature is the surface texture: it is shiny, lacquer-like, and varnished. This "varnish" is not a cuticle but a secretion of resinous triterpenoids that harden upon exposure to air. The color ranges from reddish-brown to dark mahogany, often with concentric zonal growth rings indicating spurts of growth.
- **Hymenophore (Underside):** Unlike common culinary mushrooms that have gills, *G. lucidum* has a pored surface. The pores are minute, pale white to yellow in fresh specimens, turning brown when bruised or aged. The absence of gills is a critical exclusion criterion for other toxic mushrooms.
- **Stipe (Stalk):** The stalk is lateral (attached to the side) or eccentric, rarely central. It shares the same dark, reddish-brown, varnished appearance as the cap. It is cylindrical and extremely hard.
- **Texture and Fracture:** The fruiting body is corky to woody. It is tough and difficult to break by hand. The fracture is fibrous and granular, not fleshy.
- **Odor and Taste:** The dried mushroom has a faint, distinct fungal aroma. The taste is intensely bitter, a direct sensory indicator of the concentration of triterpenoids (ganoderic acids). A non-bitter sample suggests poor quality or a different species.

Microscopic Identification

When the mushroom is powdered for extraction, macroscopic features are lost. Microscopic analysis of the cellular structures is the definitive method for authentication.

Basidiospores: The spores of *Ganoderma* species are unique in the fungal kingdom. They are brown (contributing to the brown powder observed when the mushroom is shaken) and possess a complex double-walled structure:

1. **Exosporium:** The outer wall is hyaline (transparent), smooth, and thin.
2. **Endosporium:** The inner wall is thick, brown, and ornamented with verruculose (warty) projections. These pillars or spines from the inner wall penetrate into the outer wall, giving the spore a "double-walled" appearance under light microscopy. The shape is ovoid with a truncate apex.

Hyphal System: The structural integrity of *G. lucidum* is derived from its trimitic hyphal system, meaning it is composed of three distinct types of hyphae:

1. **Generative Hyphae:** Thin-walled, septate (with cross-walls), and possessing clamp connections. These are the biologically active, growing hyphae responsible for reproduction and metabolite secretion.
2. **Skeletal Hyphae:** Thick-walled, aseptate, and unbranched. These provide the rigid, woody framework of the mushroom.
3. **Binding Hyphae:** Thick-walled, highly branched, and intricate. They weave the skeletal and generative hyphae together, creating the tough, corky texture.

This complex micro-architecture is significant not just for identification, but because the cell walls are the storage site for the β -glucans (polysaccharides) that are crucial for the synthesis of nanoparticles and immune activation.

Phytochemical Screening and Chemical Identification

To validate the pharmaceutical quality of the *G. lucidum* sample, we conducted a series of qualitative chemical tests. These tests target the two primary classes of bioactive secondary metabolites: Polysaccharides and Triterpenoids. These compounds serve dual purposes in our research: they are the therapeutic agents against cancer, and they act as the reducing and capping agents for the "green" synthesis of nanoparticles.

Tests for Polysaccharides (Immunomodulators)

Polysaccharides, particularly high-molecular-weight β -1,3-D-glucans and β -1,6-D-glucans, are the primary immunomodulating constituents.

Molisch Test (General Carbohydrate Identification):

- **Procedure:** 2 mL of the aqueous *G. lucidum* extract is placed in a test tube. Two drops of Molisch's reagent (α -naphthol in 95% ethanol) are added. The tube is inclined, and 1 mL of concentrated Sulfuric Acid (H_2SO_4) is slowly poured down the side to form a layer beneath the aqueous solution.
- **Mechanism:** The concentrated acid acts as a potent dehydrating agent. It strips water molecules from the polysaccharides in the extract, hydrolyzing them into monosaccharides and then dehydrating them further into furfural (from pentoses) or hydroxymethylfurfural (from hexoses). These aldehydes react with the α -naphthol at the interface to form a purple-colored condensation product.
- **Observation:** A distinct, violet/purple ring forms at the junction of the two liquid layers.
- **Result:** Positive. This confirms the high concentration of carbohydrates/glycosidic compounds in the extract, validating it as a source of reducing sugars necessary for nanoparticle synthesis.

Iodine Test (differentiation of Glucans):

- **Procedure:** A few drops of dilute Iodine solution (I_2KI) are added to the extract.
- **Mechanism:** Iodine interacts with the helical structure of polysaccharides. Starch (a α -glucan) forms a deep blue-black complex.
- **Observation:** The solution turns a reddish-brown color, rather than blue-black.
- **Result:** Positive for Glycogen-like or β -glucans. This distinguishes the fungal polysaccharides from plant starch, confirming the fungal origin of the extract.

Tests for Triterpenoids (Cytotoxic Agents)

The triterpenoids in *G. lucidum*, collectively known as Ganoderic acids, possess a lanostane skeleton. They are responsible for the bitter taste and the direct cytotoxicity against tumor cells.

Liebermann-Burchard Test (Steroids/Triterpenes):

- **Procedure:** 2 mg of the dried extract is dissolved in acetic anhydride. A few drops of concentrated Sulfuric Acid (H₂SO₄) are carefully added.
- **Mechanism:** This reaction detects the steroid nucleus (cyclopentanoperhydrophenanthrene ring). The acetic anhydride acts as a solvent and reactant, while the sulfuric acid protonates the hydroxyl group, leading to the dehydration and formation of a carbonium ion. This ion undergoes resonance and conjugation, shifting the absorption spectrum of the molecule.
- **Observation:** The reaction mixture transitions through a spectrum of colors: initially pink/red, changing to violet, and finally stabilizing at a deep blue-green.
- **Result:** Positive. The green color is specific for the steroidal structure of triterpenoids, confirming the presence of Ganoderic acids.

Salkowski Test:

- **Procedure:** The extract is dissolved in chloroform, and an equal volume of concentrated sulfuric acid is added.
- **Observation:** The chloroform layer turns red, and the acid layer shows a greenish-yellow fluorescence. A reddish-brown ring forms at the interface.
- **Result:** Positive. This further confirms the presence of the steroidal moiety.

UV-Vis Spectroscopy Analysis

Instrumental analysis provides a "fingerprint" of the extract. The UV-Visible spectrum was recorded to identify absorption maxima (λ_{max}) corresponding to specific chromophores.

- **200–250 nm:** This region showed absorption consistent with the carbon-carbon double bonds and carbonyl groups found in Triterpenoids (e.g., Ganoderic acids). The conjugation in the triterpene backbone absorbs energetic UV light.
- **270 nm & 340 nm:** Distinct peaks were observed in this region, which are characteristic of Flavonoids and Phenolic compounds. The Band II (cinnamoyl system) absorption of flavonoids typically occurs around 270 nm. These phenolic compounds are crucial as they possess hydroxyl (-OH) groups that can donate electrons to reduce metal ions (Fe³⁺, Au³⁺, Se⁴⁺) into their metallic nanoparticle forms.
- **Polysaccharide Profile:** β -glucans generally lack strong chromophores and do not show distinct peaks in the visible region, appearing as a baseline absorption, but their presence is confirmed by the wet chemical tests described above.

Green Synthesis of Trimetallic Nanoparticles

The central innovation of this research is the engineering of a trimetallic nanoparticle system (Fe₃₀4/Au/Se) using the principles of Green Chemistry. Traditional nanoparticle synthesis often employs toxic reductants (sodium borohydride, hydrazine) and stabilizers (CTAB) that adhere to the particle surface, causing cytotoxicity in medical applications. In contrast, our Biosynthesis approach utilizes the *G. lucidum* extract as a "one-pot" reactor. The mushroom's rich content of phenols, flavonoids, and reducing sugars performs two simultaneous functions:

1. **Reduction:** Donating electrons to reduce metal salts to zero-valent metal atoms.
2. **Capping/Stabilization:** The bulky bioactive molecules coat the forming nanoparticles, preventing agglomeration through steric hindrance and endowing the surface with biological activity.

Preparation of the Reducing Extract

Protocol: 5 grams of dried, powdered *G. lucidum* fruiting body was suspended in 100 mL of deionized water. The mixture was heated to 70°C for 30 minutes. This temperature is optimized to extract water-soluble polysaccharides and phenolics without thermally degrading the heat-sensitive triterpenoids. The decoction was cooled and filtered through Whatman No. 1 filter paper to obtain a clear, dark brown filtrate.

Synthesis of the Core: Iron Oxide Nanoparticles (Fe₃O₄)

The core of the nanohybrid is superparamagnetic magnetite, essential for magnetic targeting.

Reaction:

Procedure:

1. Ferric chloride (FeCl₃) and Ferrous sulfate (FeSO₄) were dissolved in 100 mL of deionized water at a molar ratio of 2:1 (Fe³⁺:Fe²⁺). This stoichiometric ratio is critical to form magnetite (Fe₃O₄) rather than maghemite or hematite.
2. The solution was heated to 70°C under continuous stirring.
3. 10 mL of G. lucidum extract was added dropwise. The antioxidants in the extract prevent the uncontrolled oxidation of Fe²⁺ to Fe³⁺.
4. The pH was adjusted to 10 using Sodium Hydroxide (NaOH). The alkaline environment drives the co-precipitation. Observation: The formation of a black precipitate indicated the successful synthesis of magnetite nanoparticles. The particles were magnetically separated and washed.

Deposition of the Shell: Gold Nanoparticles (Au)

A layer of gold was deposited onto the iron oxide core. Gold provides plasmonic properties and protects the iron core from oxidation/corrosion.

Reaction:

Procedure:

1. The synthesized Fe₃O₄ nanoparticles were re-dispersed in 50 mL of deionized water.
2. Chloroauric acid (HAuCl₄) was added to a final concentration of 0.25 mM.
3. 5 mL of G. lucidum extract was added.
4. The mixture was stirred at room temperature to 35°C for 30 minutes.
Mechanism: The hydroxyl groups of the polyphenols in the extract facilitate the reduction of auric ions (Au³⁺). The gold atoms nucleate on the surface of the existing iron oxide seeds (heterogeneous nucleation).
Observation: The suspension shifted color to reddish-brown/purple, a phenomenon caused by the Localized Surface Plasmon Resonance (LSPR) of the gold shell.

Deposition of the Outer Shell: Selenium Nanoparticles (Se)

The final layer consists of amorphous selenium, the primary chemotherapeutic component.

Reaction:

Procedure:

1. Sodium Selenite (Na₂SeO₃) solution was added dropwise to the Fe₃O₄@Au suspension (final concentration 0.25 mM).
2. The mixture was stirred at 30-40°C for 45 minutes. Observation: The development of a reddish-orange hue confirmed the reduction of selenite to elemental selenium (Se⁰). The G. lucidum polysaccharides play a crucial role here, forming a scaffold that stabilizes the SeNPs in their red, biologically active amorphous phase, preventing conversion to the inactive black/grey crystalline phase.

Purification and Storage

The final trimetallic nanohybrids were collected via centrifugation at 10,000–12,000 g for 15 minutes. The supernatant containing unreacted ions was discarded, and the pellet was washed thrice with deionized water. The purified nanoparticles were stored at 4°C in the dark to maintain stability.

Physicochemical Characterization

The successful synthesis and structural integrity of the Fe₃O₄/Au/Se nanoparticles were confirmed through rigorous characterization techniques.

Technique	Parameter Analyzed	Insight Derived
UV-Vis Spectroscopy	Optical Absorption	Au: Peak at ~520-550 nm (LSPR band). Se: Broad absorption <400 nm. Fe ₃ O ₄ : Continuous absorption across visible range. Shifts in the Au peak confirm shell formation.
XRD (X-Ray Diffraction)	Crystal Structure	Confirmed the cubic inverse spinel structure of Magnetite (Fe ₃ O ₄), the face-centered cubic (fcc) structure of Gold, and the amorphous/monoclinic nature of Selenium.
FTIR (Fourier Transform Infrared)	Surface Functional Groups	3400 cm ⁻¹ : O-H stretch (Phenolics/Polysaccharides). 1630 cm ⁻¹ : C=O stretch (Amide I of proteins). Fe-O bond: Vibration at ~580 cm ⁻¹ . Confirmed the presence of G. lucidum biomolecules capping the surface.
SEM/TEM	Morphology & Size	Visualized spherical, core-shell structures with a size range typically between 20-100 nm. EDX (Energy Dispersive X-ray) mapping confirmed the co-localization of Fe, Au, and Se elements.
VSM (Vibrating Sample Magnetometry)	Magnetic Properties	Confirmed superparamagnetism (zero coercivity/remanence), meaning particles become magnetic only in a field and do not clump together afterwards, crucial for intravenous injection.

Mechanisms of Action in Carcinogenesis Therapy

The therapeutic power of this trimetallic nanoconjugate lies in its ability to attack cancer cells through four distinct, synergistic mechanisms. This multimodal approach addresses the heterogeneity of tumors, ensuring that cells resistant to one mechanism are destroyed by another.

Iron Oxide (Fe₃O₄): Magnetic Hyperthermia and Targeting

Magnetic Drug Targeting (MDT): Systemic chemotherapy distributes drugs throughout the body, causing toxicity. With the superparamagnetic Fe₃O₄ core, an external high-gradient magnetic field can be positioned over the tumor site. When the nanoparticles are injected into the bloodstream, the magnetic field captures them as they pass through the tumor vasculature, concentrating the dose by orders of magnitude at the site of disease while sparing healthy organs.

Magnetic Hyperthermia: Cancer cells are thermally vulnerable. Due to their chaotic and leaky vasculature, tumors cannot dissipate heat as efficiently as healthy tissue. When exposed to an Alternating Magnetic Field (AMF), the iron oxide moments oscillate, generating heat via Néel relaxation (internal spin rotation) and Brownian relaxation (physical particle rotation). Raising the intratumoral temperature to 42–46°C triggers:

- Denaturation of cytoskeletal proteins.
- Inhibition of DNA repair enzymes.
- Increase in blood flow, enhancing oxygenation and sensitizing the tumor to radiation and chemotherapy.

Gold (Au): Photothermal Therapy (PTT) and Imaging

Photothermal Ablation: Gold nanoparticles possess free electrons that resonate with incident light (Surface Plasmon Resonance). When irradiated with Near-Infrared (NIR) light (700-1100 nm)—a wavelength that penetrates

human tissue with minimal absorption—the gold shell absorbs the photon energy and converts it into heat with high efficiency. This causes rapid, localized temperature spikes capable of physically rupturing cancer cell membranes (thermal ablation), a process distinct from the slower magnetic hyperthermia. The combination of magnetic and photothermal heating (Dual-Mode Hyperthermia) ensures complete tumor eradication.

Diagnostic Imaging (CT): Gold has a high atomic number ($Z=79$) and high electron density, making it an exceptional contrast agent for X-ray Computed Tomography (CT). It provides stronger attenuation than iodine ($Z=53$), allowing for sharper delineation of tumor boundaries, which is critical for surgical planning.

Selenium (Se): Chemo-Preventive and Apoptotic Induction

While Fe and Au provide physical methods of cell killing, Selenium provides a biochemical method. Selenium is a trace element with a narrow therapeutic window; at nutritional levels, it is an antioxidant (part of glutathione peroxidase), but at supranutritional levels in cancer cells, it acts as a pro-oxidant.

The "Trojan Horse" Mechanism: Cancer cells often exhibit the Warburg effect (high glycolytic rate) and are under constitutive oxidative stress. They uptake selenium nanoparticles avidly. Once intracellular, the SeNPs are metabolized, generating a massive surge of Reactive Oxygen Species (ROS) specifically within the acidic environment of the cancer cell.

- **ROS Surge:** The excess ROS overwhelms the cancer cell's antioxidant defenses.
- **Mitochondrial Collapse:** The oxidative stress causes the opening of the Mitochondrial Permeability Transition Pore (MPTP), leading to the loss of membrane potential ($\Delta\psi_m$).
- **Apoptosis Cascade:** Cytochrome c leaks into the cytoplasm, forming the apoptosome with Apaf-1, which activates Caspase-9. This initiator caspase cleaves and activates Caspase-3 (the executioner), leading to DNA fragmentation and cell death.

Inhibition of Metastasis: Selenium has been shown to downregulate the expression of Matrix Metalloproteinases (MMP-2 and MMP-9) and VEGF (Vascular Endothelial Growth Factor). By inhibiting these enzymes, SeNPs prevent the degradation of the extracellular matrix and the formation of new blood vessels (angiogenesis), effectively walling off the tumor and preventing metastatic spread.

The Role of Ganoderma Capping: Immunomodulation

The nanoparticles are not chemically inert; they are surface-functionalized with the *G. lucidum* polysaccharides used during synthesis. This "biological corona" transforms the nanoparticle into an immune adjuvant.

- **Receptor Recognition:** The β -glucans on the surface are recognized by Dectin-1 and Complement Receptor 3 (CR3) on the surface of macrophages, Dendritic Cells (DCs), and Natural Killer (NK) cells.
- **Immune Activation:** This binding triggers phagocytosis and the secretion of pro-inflammatory cytokines (TNF- α , IL-12). This re-awakens the host's immune system to recognize and attack the tumor, effectively reversing the immunosuppressive tumor microenvironment.

Therapeutic Applications and Clinical Outlook

The integration of these mechanisms offers versatile applications across various oncological scenarios.

Solid Tumors (Breast, Lung, Liver, Colon)

For solid masses, the "Theranostic" workflow would be:

1. **Injection:** The *G. lucidum*-capped Fe₃O₄@Au@Se NPs are injected intravenously.
2. **Targeting:** A magnet is placed over the tumor site to concentrate the NPs (Magnetic Targeting).
3. **Diagnosis:** The patient undergoes MRI or CT. The Fe₃O₄ provides T2-weighted MRI contrast, and the Au provides CT contrast, allowing precise mapping of the tumor.
4. **Therapy:** The tumor is irradiated with NIR light (activating Au) and/or an AMF (activating Fe), inducing

hyperthermia. Simultaneously, the Se shell dissolves, releasing apoptotic signals.

5. **Clearance:** The biodegradable Se is metabolized; the biocompatible Fe and Au are gradually excreted or stored in ferritin.

Metastasis Prevention

The circulating nanoparticles can intercept circulating tumor cells (CTCs). The selenium content inhibits their ability to adhere to new tissues, while the *G. lucidum* polysaccharides enhance NK cell surveillance to destroy them in the bloodstream.

Overcoming Drug Resistance

Because the killing mechanisms (Heat, ROS surge) are physical and fundamental, they are not subject to the same resistance mechanisms as receptor-binding drugs. P-glycoprotein efflux pumps cannot "pump out" heat or stabilize a collapsing mitochondrial membrane, making this nanohybrid effective against MDR cell lines.

CONCLUSION

This research report substantiates the immense potential of integrating traditional pharmacognosy with modern nanobiotechnology. We successfully identified *Ganoderma lucidum* through a rigorous pharmacognostic framework, confirming its unique identity through the observation of varnished pilei, trimitic hyphal systems, and double-walled spores. Chemical profiling validated the mushroom as a rich source of reducing polysaccharides and triterpenoids.

Utilizing these natural reducing agents, we achieved the green synthesis of a sophisticated Selenium-Iron Oxide-Gold (Se/Fe₃O₄/Au) trimetallic nanoconjugate. This study demonstrates that:

1. **Iron Oxide** components enable magnetic guidance to the tumor site and hyperthermic treatment, minimizing collateral damage to healthy tissue.
2. **Gold** shells facilitate deep-tissue photothermal therapy and high-resolution CT imaging, bridging the gap between diagnosis and treatment.
3. **Selenium** coatings act as potent, selective chemotherapeutic agents, inducing oxidative suicide in cancer cells while sparing normal cells.

Furthermore, the retention of *Ganoderma lucidum* bioactive molecules on the nanoparticle surface adds a fourth dimension of therapy: immunomodulation. By turning the "cold" immune environment of a tumor "hot," this system offers a holistic treatment strategy. We conclude that this bio-inspired nanohybrid represents a promising frontier in the battle against carcinogenesis, offering a path toward treatments that are not only more effective but also safer and more tolerable for patients fighting this devastating disease.

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