**Smart Nanomaterials in Drug Delivery System**

**Author**

Prakash Kumar Sahoo

Post Graduate Department of Chemistry

Dhenkanal Autonomous College

Dhenkanal, Odisha-759001

Correspondence Email- prakashchemistry1989@gmail.com

**Abstract**

The field of drug delivery has witnessed a paradigm shift with the advent of smart nanomaterials. These advanced materials, designed at the nanoscale, offer unprecedented control over drug release, targeting, and therapeutic efficacy. This abstract provides an overview of the transformative potential of smart nanomaterials in drug delivery systems. Smart nanomaterials are engineered to respond to specific stimuli, such as changes in pH, temperature, or enzymatic activity, allowing for precise and controlled drug release at the desired site of action. This targeted drug delivery minimizes off-target effects, reduces side effects, and enhances the therapeutic index of drugs, thereby improving patient outcomes. Furthermore, smart nanomaterials can encapsulate a wide range of therapeutics, including small molecules, proteins, nucleic acids, and even gene-editing tools. This versatility makes them a promising platform for the treatment of various diseases, including cancer, infectious diseases, neurodegenerative disorders, and more.

In addition to controlled drug release, smart nanomaterials enable real-time monitoring and feedback through integrated sensors or imaging agents. This capability allows for personalized medicine approaches, where treatment regimens can be adjusted in response to individual patient needs. This abstract also discusses the challenges and considerations in the development of smart nanomaterials for drug delivery, including safety, scalability, and regulatory hurdles. It highlights recent advancements in the field, such as the use of biocompatible and biodegradable materials, as well as the incorporation of artificial intelligence and machine learning for predictive modeling of drug release kinetics. Smart nanomaterials represent a transformative approach to drug delivery, offering precise control, enhanced therapeutic outcomes, and the potential for personalized medicine. As research in this field continues to progress, smart nanomaterials hold great promise for revolutionizing the way we administer and benefit from therapeutic agents in healthcare.

**Keywords**: nanomaterials, enzymatic activity, drug delivery, biocompatible, biodegradable etc.

**Introduction**

The advent of smart nanomaterials has ushered in a new era in the field of drug delivery, promising groundbreaking advancements in the way we administer therapeutic agents to combat diseases. Traditional drug delivery systems have often been limited by issues of off-target effects, low drug solubility, rapid clearance from the body, and the inability to precisely control drug release. In response to these challenges, smart nanomaterials have emerged as a revolutionary solution, offering precise control over drug delivery, improved targeting, and enhanced therapeutic outcomes. Smart nanomaterials are engineered at the nano scale, allowing for a level of precision that was previously unattainable in drug delivery. These nano systems are designed to respond to specific stimuli, which can include changes in pH, temperature, enzymatic activity, or external triggers like light or magnetic fields. Such responsiveness enables these nanomaterials to release therapeutic agents in a highly controlled and site-specific manner, mitigating the risk of adverse side effects and enhancing the overall therapeutic index of drugs.

The potential applications of smart nanomaterials in drug delivery are vast, encompassing a wide range of therapeutic agents, from small molecules to biologics such as proteins, nucleic acids, and even emerging gene-editing tools. This versatility opens the door to treating a myriad of diseases, including cancer, infectious diseases, neurodegenerative disorders, and more. One of the key advantages of smart nanomaterials lies in their ability to provide real-time monitoring and feedback. Incorporating sensors or imaging agents within these nano systems allows for the precise tracking of drug release, tissue distribution, and therapeutic efficacy. This capability paves the way for personalized medicine approaches, where treatment regimens can be tailored to individual patient needs, maximizing the chances of successful outcomes while minimizing adverse effects.

Despite the immense promise of smart nanomaterials, their development is not without challenges. Issues related to safety, scalability, and regulatory approvals must be carefully addressed. Researchers are continuously exploring innovative strategies, including the use of biocompatible and biodegradable materials, to overcome these hurdles and bring smart nano material based drug delivery systems closer to clinical reality. In this comprehensive exploration of smart nanomaterials for drug delivery, we will delve into the design principles, applications, recent advancements, and future prospects of these groundbreaking technologies. By harnessing the power of nano scale engineering and responsive materials, we have the potential to transform the landscape of healthcare, offering patients more effective and personalized treatment options than ever before.

**Design rationale of smart drug delivery nanoplatforms**

The design rationale of smart drug delivery nano platforms is driven by the need to overcome various challenges associated with conventional drug delivery systems. These challenges include limited drug solubility, poor bioavailability, off-target effects, rapid drug clearance, and the inability to provide precise control over drug release. Smart drug delivery nano platforms are designed to address these issues while offering several key advantages:

1. **Precise Drug Targeting :**

Smart nano platforms are engineered to deliver drugs to specific target sites, such as tumors or diseased tissues, with high precision. This targeting minimizes damage to healthy cells and tissues, reducing side effects.

**(ii) Controlled Drug Release:** These nano systems respond to specific stimuli or triggers, allowing for controlled drug release at the desired location and time. Common triggers include changes in pH, temperature, enzymatic activity, or external factors like light or magnetic fields. This control optimizes therapeutic efficacy and minimizes fluctuations in drugconcentration.

**(iii) Enhanced Drug Stability**: Nano carriers can protect drugs from degradation, improving their stability and shelf life. This is especially important for sensitive drugs, such as proteins and nucleic acids.

**(iv) Improved Drug Solubility**: Many drugs suffer from poor solubility in physiological fluids. Nano platforms can encapsulate hydrophobic drugs, improving their solubility and bioavailability.

**(v) Reduced Immunogenicity**: Smart nanomaterials can be designed to be biocompatible and biodegradable, minimizing the risk of triggering immune responses when delivering biotherapeutics.

**(vi) Personalized Medicine**: The integration of sensors and imaging agents into nano platforms allows for real-time monitoring of drug release and therapeutic responses. This information can be used to customize treatment regimens for individual patients, maximizing effectiveness.

**(vii) Minimized Drug Resistance**: Targeted drug delivery can help reduce the development of drug resistance in diseases like cancer by concentrating therapeutic agents at the site of action.

The design process of smart drug delivery nano platforms typically involves the following steps:

**(Step-I) Material Selection**: Choosing appropriate nanomaterials that are biocompatible, biodegradable, and capable of responding to the desired stimuli. Common materials include lipids, polymers, and inorganic nanoparticles.

**(Step-II) Drug Loading**: Efficient loading of the drug into the nanocarrier while maintaining its stability and bioactivity.

**(Step-III) Stimuli-Responsive Elements**: Incorporating stimuli-responsive elements, such as pH-sensitive polymers or thermo-responsive materials, to enable controlled drug release.

**(Step-IV) Surface Functionalization**: Modifying the surface of nano carriers with targeting ligands, antibodies, or peptides to enhance their ability to bind to specific cells or tissues.

**(Step-V) Safety and Toxicity Assessment**: Evaluating the safety and potential toxicity of the nano platform components through in vitro and in vivo studies.

**(Step-VI) In Vitro and In Vivo Testing**: Testing the smart nano platform in cell cultures and animal models to assess its performance, targeting efficiency, and therapeutic efficacy.

**(Step-VII) Regulatory Considerations**: Addressing regulatory requirements and ensuring compliance with safety and efficacy standards for clinical translation.

The design rationale of smart drug delivery nano platforms is guided by the goal of optimizing drug delivery while minimizing side effects and maximizing therapeutic benefits. As research in nanotechnology and materials science continues to advance, these platforms hold significant promise for the development of innovative and personalized treatment strategies across various medical disciplines.

1. **pH-responsive Smart Nanomaterials in drug delivery**

pH-responsive smart nanomaterials play a pivotal role in drug delivery systems by providing a means to control drug release in response to changes in the local pH environment. These materials are designed to respond to variations in pH, typically between the acidic environment of specific diseased tissues (e.g., tumor microenvironments) and the near-neutral pH of healthy tissues. This selective drug release enables precise targeting and improved therapeutic outcomes. Here's an overview of pH-responsive smart nanomaterials in drug delivery:

**1. Mechanism of pH Responsiveness**:

**pH-sensitive Polymers**: Many pH-responsive nanomaterials are constructed using polymers with pH-sensitive functional groups, such as carboxylic acids or amines. These polymers can undergo conformational changes or protonation/deprotonation in response to pH fluctuations, leading to alterations in the nanomaterial's structure and drug release properties.

**Surface Charge Reversal**: pH-responsive nanocarriers can change their surface charge in response to pH changes. For example, at low pH (acidic) conditions, nanocarriers with pH-responsive coatings may become positively charged, facilitating cellular uptake in acidic environments, such as those found in tumor tissues.

**2. Advantages of pH-Responsive Nanomaterials**:

**(i) Enhanced Targeting**: pH-responsive nanomaterials can take advantage of the acidic pH found in various disease microenvironments, including tumors and inflamed tissues. This property allows for selective drug release at disease sites, minimizing off-target effects.

**(ii) Controlled Drug Release**: By designing nanomaterials that respond to specific pH ranges, drug release can be precisely controlled, ensuring that therapeutic agents are released when and where they are needed most.

**(iii) Improved Therapeutic Efficacy**: The ability to maintain therapeutic drug concentrations within the target tissue for longer durations can enhance the overall efficacy of treatments.

**(iv) Reduced Side Effects**: Selective drug release minimizes exposure to healthy tissues, reducing the risk of side effects commonly associated with systemic drug administration.

**3. pH-Responsive Nanomaterial Types**:

**(i) Liposome**: pH-responsive liposome are lipid-based nano carriers that can encapsulate both hydrophilic and hydrophobic drugs. They can be engineered to release their cargo in response to acidic conditions, making them suitable for targeting tumours and intracellular drug delivery.

**(ii) Polymeric Nanoparticles**: pH-responsive polymer nanoparticles can be designed to disintegrate or swell under acidic conditions, leading to drug release. These nanoparticles offer versatility in terms of drug loading and delivery applications.

**(iii) Micelles**: pH-responsive micelles are self-assembling structures formed by amphiphilic block copolymers. They can encapsulate hydrophobic drugs and release them in response to pH changes, making them useful for intracellular drug delivery.

**(iv) Hydrogels**: pH-responsive hydrogels can be injected or implanted at specific sites. They can release drugs in response to pH changes, making them suitable for localized drug delivery, wound healing, or tissue engineering applications.

**4. Clinical Applications**:

pH-responsive smart nanomaterials have shown promise in the treatment of cancer, where the tumor microenvironment is typically acidic. They are also explored for inflammatory diseases, infections, and conditions involving pH variations in specific tissues. So pH-responsive smart nanomaterials represent a powerful and versatile tool in drug delivery systems. Their ability to precisely control drug release in response to pH changes allows for improved targeting, reduced side effects, and enhanced therapeutic efficacy in various clinical applications. Researchers continue to refine and expand the use of pH-responsive nanomaterials to advance the field of personalized medicine and tailored drug delivery.Top of Form

1. **Redox-responsive Smart Nanomaterials in drug delivery**

Redox-responsive smart nanomaterials are a class of advanced drug delivery systems designed to release therapeutic agents in response to changes in the redox (reduction-oxidation) environment within cells or tissues. The redox potential in biological systems can vary significantly between healthy and diseased tissues, making redox-responsive nanomaterials a promising strategy for targeted drug delivery. Here's an overview of redox-responsive smart nanomaterials in drug delivery:

1. **Mechanism of Redox Responsiveness**:

**(i) Redox-Active Components**: Redox-responsive nanomaterials typically incorporate redox-active moieties or bonds, such as disulfide (S-S) linkages, thioether (S-C) bonds, or selenide (Se) groups, within their structure. These components can be cleaved or altered in response to variations in the intracellular redox environment.

**(ii) Glutathione (GSH) Sensing**: GSH is a crucial intracellular antioxidant with high reducing potential. Redox-responsive nanomaterials often exploit the presence of GSH within cells. In a reducing environment (high GSH concentration), the redox-active bonds are cleaved, leading to the release of the encapsulated drug.

**2. Advantages of Redox-Responsive Nanomaterials**:

**(i) Intracellular Drug Delivery**: Redox-responsive systems are designed to respond to the reducing conditions found within cells, enabling targeted drug release inside the desired cellular compartments.

**(ii) Minimized Off-Target Effects**: The ability to release drugs specifically within cells or tissues with elevated levels of GSH reduces the risk of off-target effects and systemic exposure.

**(iii) Enhanced Drug Stability**: By encapsulating drugs within redox-responsive nanomaterials, the stability and bioavailability of certain drugs can be improved, especially those susceptible to degradation in the extracellular environment.

**(iv) Increased Therapeutic Efficacy**: The controlled release of drugs within cells or specific tissues can enhance their therapeutic efficacy by maintaining therapeutic concentrations over an extended period.

**3. Redox-Responsive Nanomaterial Types**:

**(i) Polymeric Nanoparticles**: Nanoparticles constructed from redox-responsive polymers, often containing disulfide bonds, can disintegrate in response to the intracellular redox environment, leading to drug release.

**(ii) Liposomes**: Redox-responsive liposomes can be designed to contain disulfide bonds in their lipid bilayers, allowing for the release of encapsulated drugs in response to intracellular GSH.

**(iii) Micelles**: Redox-responsive micelles, formed from amphiphilic block copolymers containing redox-sensitive linkages, can encapsulate hydrophobic drugs and disassemble under reducing conditions.

**4. Clinical Applications**:

Redox-responsive smart nanomaterials hold significant potential for the treatment of various diseases, including cancer. Tumor cells often exhibit higher levels of GSH compared to normal cells, making them prime targets for redox-responsive drug delivery. These nanomaterials can also be employed in the treatment of neurodegenerative diseases, inflammatory conditions, and infections, where intracellular delivery or targeted drug release is critical.

**5. Challenges and Considerations**:

(i) Ensuring the biocompatibility and safety of redox-responsive nanomaterials is essential for their clinical translation.

(ii) The complexity of intracellular redox environments may require fine-tuning of nanomaterials to achieve precise drug release.

(iii) Regulatory considerations and long-term stability of these nanomaterials need to be addressed for clinical applications.

Lastly, redox-responsive smart nanomaterials offer a promising avenue for targeted drug delivery, particularly within intracellular compartments where redox conditions vary. These nanomaterials have the potential to enhance the therapeutic efficacy of drugs while minimizing off-target effects, advancing the field of personalized medicine and tailored drug delivery.Top of Form

1. **Temperature-responsive Smart Nanomaterials in drug delivery**

Temperature-responsive smart nanomaterials are a class of advanced drug delivery systems designed to release therapeutic agents in response to changes in temperature. These materials exhibit reversible phase transitions within a specific temperature range, which can be exploited for precise drug release and enhanced therapeutic outcomes. Here's an overview of temperature-responsive smart nanomaterials in drug delivery:

**1. Mechanism of Temperature Responsiveness**:

**(i) Thermoresponsive Polymers**: Temperature-responsive nanomaterials are often constructed using thermoresponsive polymers, such as poly(N-isopropylacrylamide) (PNIPAM). These polymers undergo a phase transition, typically around their lower critical solution temperature (LCST), where they change from a hydrophilic to a hydrophobic state.

**(ii) Solubility Changes:** Below the LCST, the polymer is in a hydrophilic state, allowing for drug encapsulation and stable nanomaterial formation. Above the LCST, the polymer becomes hydrophobic, leading to nanomaterial disassembly and drug release.

**2. Advantages of Temperature-Responsive Nanomaterials:**

**(i) Precise Control**: Temperature-responsive nanomaterials offer precise control over drug release, triggered by external temperature changes or the inherent temperature variations within the body.

**(ii) Minimized Off-Target Effects**: The ability to release drugs in a controlled manner at a specific temperature range reduces the risk of off-target effects and allows for site-specific drug delivery.

**(iii) Enhanced Drug Stability**: By encapsulating drugs within temperature-responsive nanomaterials, the stability and solubility of certain drugs can be improved, especially those susceptible to temperature-induced degradation.

**(iv) Local Hyperthermia**: In some cases, temperature-responsive nanomaterials can be used in combination with local hyperthermia techniques, where heat is applied externally to increase the temperature within the target tissue. This can further enhance drug release and therapeutic effects.

**3. Temperature-Responsive Nanomaterial Types:**

**(i) Liposome**: Temperature-responsive liposomes can be designed to contain thermo-sensitive lipids or polymers. Above the LCST, these liposomes undergo phase transition, leading to drug release.

**(ii) Polymeric Nanoparticles:** Nanoparticles constructed from temperature-responsive polymers, including PNIPAM, can be tailored to disassemble at specific temperatures, facilitating drug release.

**(iii) Hydrogels**:Temperature-responsive hydrogels can be used for localized drug delivery. They can be injected or implanted and respond to temperature changes, releasing drugs within the target tissue.

**4. Clinical Applications:**

Temperature-responsive smart nanomaterials have potential applications in various diseases, including cancer. External heating techniques, such as hyperthermia, can be used to trigger drug release within tumors. These nanomaterials can also be employed in the treatment of inflammatory conditions, infections, and tissue engineering, where precise control over drug release is desired.

**5. Challenges and Considerations:**

(i) Ensuring the biocompatibility and safety of temperature-responsive nanomaterials is crucial for clinical use.

(ii) Fine-tuning the LCST of these nanomaterials to match the desired therapeutic temperature range is essential for precise drug release.

(iii) External heating methods for triggering drug release may require specialized equipment and careful monitoring.

Hence, the temperature-responsive smart nanomaterials offer a versatile and controlled approach to drug delivery. Their ability to respond to temperature changes allows for precise drug release, making them valuable tools for enhancing therapeutic outcomes in various clinical applications.

1. **Light, magnetic, and US responsive Smart Nanomaterials in drug delivery**

Light, magnetic, and ultrasound (US) responsive smart nanomaterials are innovative drug delivery systems that leverage external stimuli to trigger controlled drug release. These responsive materials offer precise spatiotemporal control over drug delivery, enhancing therapeutic efficacy while minimizing off-target effects. Here's an overview of each of these responsive nanomaterials in drug delivery:

**1. Light-Responsive Smart Nanomaterials**:

Mechanism: Light-responsive nanomaterials are designed to release drugs upon exposure to specific wavelengths of light. These materials often incorporate light-absorbing molecules or photoresponsive groups that undergo conformational changes or photothermal effects when irradiated with light.

**Advantages:**

(i) **Precise Spatial Control**: Light can be precisely focused on the target site, allowing for localized drug release.

(ii) **On-Demand Release**: Drug release can be initiated or halted by controlling the duration and intensity of light exposure.

(iii) **Minimized Off-Target Effects**: Selective drug release at the target site reduces the risk of side effects.

**Applications**:

Light-responsive nanomaterials find applications in the treatment of cancer, where photodynamic therapy (PDT) can be combined with drug delivery for enhanced therapeutic outcomes. They are also explored in other diseases, such as ocular disorders, infections, and neurodegenerative conditions.

**2. Magnetic-Responsive Smart Nanomaterials**:

Mechanism: Magnetic-responsive nanomaterials are designed to respond to external magnetic fields. These materials may contain magnetic nanoparticles that can be manipulated by an external magnetic field to trigger drug release.

**Advantages:**

(i) **Remote Control**: Drug release can be remotely controlled by applying or adjusting the magnetic field.

(ii) **Enhanced Targeting**: Magnetic nanocarriers can be guided to specific target sites using external magnets.

**(iii) Potential for Combination Therapy**: Magnetic hyperthermia can be used alongside drug delivery for synergistic therapeutic effects.

**Applications**:

Magnetic-responsive nanomaterials are explored for cancer therapy, where targeted drug delivery to tumors is facilitated by external magnetic fields. They can also be used in cardiovascular diseases, neurodegenerative disorders, and other conditions where precise drug delivery is essential.

**3. Ultrasound (US)-Responsive Smart Nanomaterials**:

**Mechanism**:

US-responsive nanomaterials utilize the energy of ultrasound waves to trigger drug release. These materials can respond to changes in acoustic pressure or temperature induced by US waves.

**Advantages**:

(i) **Non-Invasive**: US waves can penetrate tissues non-invasively, allowing for targeted drug release deep within the body.

(ii) **On-Demand Release**: Drug release can be initiated or modulated by adjusting the US parameters.

(iii) **Enhanced Penetration**: US-responsive nanomaterials can improve drug penetration into tissues or enhance cellular uptake.

**Applications**:

US-responsive nanomaterials are used in various medical fields, including cancer therapy, where they can enhance the delivery of chemotherapeutic agents to tumors. They are also employed in drug delivery to the central nervous system, wound healing, and regenerative medicine.

**Challenges and Considerations**

(i) Ensuring the biocompatibility and safety of these smart nanomaterials is essential for clinical translation.

(ii) Fine-tuning the responsiveness and specificity of these materials to external stimuli requires careful design and characterization.

(iii) Regulatory approvals and scalability must be considered when developing these nanomaterials for clinical use.

Hence, the light, magnetic, and US-responsive smart nanomaterials offer versatile strategies for targeted drug delivery. Their ability to respond to external stimuli allows for precise control over drug release, making them valuable tools for enhancing therapeutic outcomes in various medical applications.

1. **Smart nanoscale Drug Delivery System**

A smart nanoscale drug delivery system refers to an advanced approach in the field of drug delivery that utilizes nanoscale materials and technologies to enhance the precision, control, and effectiveness of drug administration. These systems are designed to respond to specific stimuli or conditions, allowing for targeted drug release and improved therapeutic outcomes. Here are key aspects of smart nanoscale drug delivery systems:

**1. Nanoscale Materials:**

**(i) Nanoparticles**: These are tiny particles with dimensions in the nanometer range. Various materials, including polymers, lipids, and inorganic substances, can be used to create nanoparticles for drug delivery.

**(ii) Nanocarriers**: Nanocarriers are nanoscale vehicles designed to encapsulate and transport drugs. They can protect drugs from degradation, improve solubility, and enable controlled release.

**2. Stimulus-Responsive Properties:**

**(i) Stimuli**: Smart nanoscale drug delivery systems are responsive to specific stimuli, such as changes in pH, temperature, enzymatic activity, light, magnetic fields, or ultrasound. These stimuli trigger drug release or alter the properties of the nanomaterials.

**(ii) Controlled Release**:The responsiveness to stimuli enables precise control over drug release, ensuring that therapeutic agents are delivered at the desired location and time.

**3. Targeted Drug Delivery:**

**(i) Active Targeting**: Smart nanoscale systems can be engineered to actively target specific cells, tissues, or organs by incorporating ligands, antibodies, or peptides that bind to receptors on the target cells.

**(ii) Passive Targeting:** These systems can also passively target disease sites, such as tumors, by taking advantage of characteristics like enhanced permeability and retention (EPR) effects in tumor vasculature.

**4. Advantages:**

**(i) Minimized Side Effects**: Targeted drug delivery minimizes exposure of healthy tissues to the therapeutic agent, reducing side effects.

**(ii) Improved Bioavailability:** Nanoscale carriers can improve the solubility and bioavailability of poorly water-soluble drugs.

**(iii) Personalized Medicine**: Some smart nanoscale systems allow for personalized medicine by monitoring drug release and adjusting treatment regimens based on individual patient responses.

**5. Clinical Applications:**

**(i) Cancer Therapy**: Smart nanoscale drug delivery systems are extensively studied for cancer treatment due to their ability to target tumors and enhance the delivery of chemotherapeutic agents.

**(ii) Infectious Diseases**: They can be used to target pathogens, enabling the controlled release of antimicrobial agents.

**(iii) Neurological Disorders**: These systems can facilitate drug delivery across the blood-brain barrier for the treatment of neurodegenerative diseases.

**6. Challenges:**

**(i) Biocompatibility and Safety**: Ensuring that the nanomaterials used in these systems are biocompatible and safe for clinical use is a significant challenge.

**(ii) Regulatory Hurdles**: Meeting regulatory requirements for approval and scaling up production can be complex.

**(iii) Clinical Translation:** Bridging the gap between laboratory research and clinical implementation requires extensive preclinical and clinical testing.

Lastly, the smart nanoscale drug delivery systems represent a promising approach to improving the precision and effectiveness of drug therapies. Their ability to respond to specific stimuli and deliver drugs with precision holds great potential for advancing medical treatments across various diseases and conditions.

**Obstacles for smart nanoplatform in potential clinical applications**

While smart nano-platforms hold significant promise in drug delivery and potential clinical applications, several obstacles and challenges need to be addressed before they can be widely adopted in clinical settings. These obstacles include:

**(i) Biocompatibility and Safety**: Ensuring that the nanomaterials used in smart nanoplatforms are biocompatible and safe for human use is a critical concern. Any potential toxicity or adverse effects on the human body must be thoroughly evaluated through preclinical and clinical studies.

**(ii) Regulatory Approvals**: Meeting regulatory requirements for the approval of smart nanoplatforms as medical devices or drug delivery systems can be a complex and lengthy process. Ensuring compliance with regulations set by health authorities is crucial but can be challenging due to the unique nature of nanomaterials.

**(iii) Scale-Up and Manufacturing**: Transitioning from laboratory-scale production to large-scale manufacturing of nanoplatforms for clinical use can be challenging. Consistency, quality control, and cost-effectiveness must be ensured during scale-up.

**(iv) Stability and Shelf Life**: Maintaining the stability of nanoplatforms over time is crucial to ensure that the drugs or therapeutic agents they carry remain effective. Stability issues can arise due to factors like aggregation, degradation, or changes in environmental conditions.

**(v) Targeting Efficacy**: Achieving precise targeting of smart nanoplatforms to the desired cells or tissues can be complex. Issues related to limited ligand-receptor interactions or variability in target expression among patients can affect targeting efficacy.

**(vi) Drug Loading and Release Control**: Ensuring efficient drug loading into nanoplatforms and precise control over drug release kinetics are critical. The ability to maintain the drug's bioactivity and therapeutic efficacy is essential.

**(vii) Immunogenicity and Clearance**: Nanomaterials may trigger immune responses, potentially leading to clearance from the body. Strategies to minimize immunogenicity and prolong circulation times need to be developed.

**(viii) Cost-Effectiveness**: Developing and producing smart nanoplatforms can be expensive. For widespread clinical adoption, cost-effectiveness and affordability must be considered.

**(ix) Patient Variability**: Patient-specific factors, such as genetics, physiology, and disease stage, can influence the performance and effectiveness of smart nanoplatforms. Personalized approaches may be necessary to optimize treatment outcomes.

**(x) Long-Term Safety**: The long-term safety of smart nanoplatforms, including potential accumulation in organs or tissues and chronic effects, must be thoroughly investigated.

**(xi) Interdisciplinary Collaboration**: Effective translation of smart nanoplatforms from research to clinical practice often requires collaboration between experts from various fields, including chemistry, materials science, pharmacology, and clinical medicine.

Despite these obstacles, ongoing research and development efforts are making significant strides in addressing these challenges. As more knowledge and experience are gained in the field, smart nanoplatforms have the potential to revolutionize drug delivery and improve the treatment of various diseases in the future.

**Conclusion**

The smart nanomaterials in drug delivery represent a transformative and promising approach to revolutionizing the field of therapeutics. These advanced materials, engineered at the nano scale, offer precise control, enhanced targeting, and improved therapeutic outcomes. Throughout this discussion, we have highlighted key aspects of smart nanomaterials in drug delivery Precise Drug Delivery, Stimuli-Responsive Properties, Enhanced Drug Stability, Improved Targeting , Personalized Medicine etc. Despite these challenges, ongoing research and development efforts continue to push the boundaries of what is possible with smart nanomaterials in drug delivery. These materials hold immense potential for the treatment of a wide range of diseases, including cancer, infectious diseases, neurodegenerative disorders, and more. As we look to the future, the continued innovation and interdisciplinary collaboration in the fields of nanotechnology, materials science, pharmacology, and clinical medicine will be key in unlocking the full potential of smart nanomaterials in drug delivery. Ultimately, these advancements have the potential to offer patients more effective and personalized treatment options while minimizing adverse effects, marking a significant leap forward in the landscape of healthcare and therapeutics.

 **References**

1. Hrubý M, Filippov SK, Štěpánek P. Smart polymers in drug delivery systems on crossroads: Which way deserves following? *European Polymer Journal.*2015;65:82–97. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=European+Polymer+Journal&title=Smart+polymers+in+drug+delivery+systems+on+crossroads:+Which+way+deserves+following?&author=M+Hrub%C3%BD&author=SK+Filippov&author=P+%C5%A0t%C4%9Bp%C3%A1nek&volume=65&publication_year=2015&pages=82-97&)]

2. Kopeček J, Yang J. Hydrogels as smart biomaterials. *Polymer International.*2007;56:1078–98. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Polymer+International&title=Hydrogels+as+smart+biomaterials&author=J+Kope%C4%8Dek&author=J+Yang&volume=56&publication_year=2007&pages=1078-98&)]

3. Lee BK, Yun YH, Park K. Smart nanoparticles for drug delivery: Boundaries and opportunities. *Chemical Engineering Science.*2015;125:158–64. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4322781/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/25684780)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Chemical+Engineering+Science&title=Smart+nanoparticles+for+drug+delivery:+Boundaries+and+opportunities&author=BK+Lee&author=YH+Yun&author=K+Park&volume=125&publication_year=2015&pages=158-64&pmid=25684780&)]

4. Bamrungsap S, Zhao Z, Chen T, Wang L, Li C, Fu T. et al. Nanotechnology in therapeutics: a focus on nanoparticles as a drug delivery system. *Nanomedicine.*2012;7:1253–71. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22931450)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nanomedicine&title=Nanotechnology+in+therapeutics:+a+focus+on+nanoparticles+as+a+drug+delivery+system&author=S+Bamrungsap&author=Z+Zhao&author=T+Chen&author=L+Wang&author=C+Li&volume=7&publication_year=2012&pages=1253-71&pmid=22931450&)]

5. Allen TM, Cullis PR. Liposomal drug delivery systems: from concept to clinical applications. *Advanced Drug Delivery Reviews.*2013;65:36–48. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23036225)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Drug+Delivery+Reviews&title=Liposomal+drug+delivery+systems:+from+concept+to+clinical+applications&author=TM+Allen&author=PR+Cullis&volume=65&publication_year=2013&pages=36-48&pmid=23036225&)]

6. Couvreur P. Nanoparticles in drug delivery: past, present and future. *Advanced Drug Delivery Reviews.*2013;65:21–3. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22580334)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Drug+Delivery+Reviews&title=Nanoparticles+in+drug+delivery:+past,+present+and+future&author=P+Couvreur&volume=65&publication_year=2013&pages=21-3&pmid=22580334&)]

7. Alvarez-Lorenzo C, Concheiro A. Smart drug delivery systems: from fundamentals to the clinic. *Chemical Communications.*2014;50:7743–65. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/24805962)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Chemical+Communications&title=Smart+drug+delivery+systems:+from+fundamentals+to+the+clinic&author=C+Alvarez-Lorenzo&author=A+Concheiro&volume=50&publication_year=2014&pages=7743-65&pmid=24805962&)]

8. Crommelin DJ, Florence AT. Towards more effective advanced drug delivery systems. *International Journal of Pharmaceutics.*2013;454:496–511. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23415662)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=International+Journal+of+Pharmaceutics&title=Towards+more+effective+advanced+drug+delivery+systems&author=DJ+Crommelin&author=AT+Florence&volume=454&publication_year=2013&pages=496-511&pmid=23415662&)]

9. Holzapfel BM, Reichert JC, Schantz J-T, Gbureck U, Rackwitz L, Nöth U. et al. How smart do biomaterials need to be? A translational science and clinical point of view. *Advanced Drug Delivery Reviews.*2013;65:581–603. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22820527)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Drug+Delivery+Reviews&title=How+smart+do+biomaterials+need+to+be?+A+translational+science+and+clinical+point+of+view&author=BM+Holzapfel&author=JC+Reichert&author=J-T+Schantz&author=U+Gbureck&author=L+Rackwitz&volume=65&publication_year=2013&pages=581-603&pmid=22820527&)]

10. Grund S, Bauer M, Fischer D. Polymers in drug delivery—state of the art and future trends. *Advanced Engineering Materials.*2011;13:B61–B87. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Engineering+Materials&title=Polymers+in+drug+delivery%E2%80%94state+of+the+art+and+future+trends&author=S+Grund&author=M+Bauer&author=D+Fischer&volume=13&publication_year=2011&pages=B61-B87&)]

11. Annabi N, Tamayol A, Uquillas JA, Akbari M, Bertassoni LE, Cha C. et al. 25th anniversary article: rational design and applications of hydrogels in regenerative medicine. *Advanced Materials.*2014;26:85–124. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3925010/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/24741694)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Materials&title=25th+anniversary+article:+rational+design+and+applications+of+hydrogels+in+regenerative+medicine&author=N+Annabi&author=A+Tamayol&author=JA+Uquillas&author=M+Akbari&author=LE+Bertassoni&volume=26&publication_year=2014&pages=85-124&pmid=24741694&)]

12. Chang H-I, Yeh M-K. Clinical development of liposome-based drugs: formulation, characterization, and therapeutic efficacy. *International Journal of Nanomedicine.*2012;7:49–60. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3260950/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22275822)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=International+Journal+of+Nanomedicine&title=Clinical+development+of+liposome-based+drugs:+formulation,+characterization,+and+therapeutic+efficacy&author=H-I+Chang&author=M-K+Yeh&volume=7&publication_year=2012&pages=49-60&pmid=22275822&)]

13. Kumari A, Yadav SK, Yadav SC. Biodegradable polymeric nanoparticles based drug delivery systems. *Colloids and Surfaces B: Biointerfaces.*2010;75:1–18. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/19782542)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Colloids+and+Surfaces+B:+Biointerfaces&title=Biodegradable+polymeric+nanoparticles+based+drug+delivery+systems&author=A+Kumari&author=SK+Yadav&author=SC+Yadav&volume=75&publication_year=2010&pages=1-18&pmid=19782542&)]

14. Rossi F, Ferrari R, Castiglione F, Mele A, Perale G, Moscatelli D. Polymer hydrogel functionalized with biodegradable nanoparticles as composite system for controlled drug delivery. *Nanotechnology.*2014;26:015602. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/25490351)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nanotechnology&title=Polymer+hydrogel+functionalized+with+biodegradable+nanoparticles+as+composite+system+for+controlled+drug+delivery&author=F+Rossi&author=R+Ferrari&author=F+Castiglione&author=A+Mele&author=G+Perale&volume=26&publication_year=2014&pages=015602&pmid=25490351&)]

15. Shimoni O, Postma A, Yan Y, Scott AM, Heath JK, Nice EC. et al. Macromolecule functionalization of disulfide-bonded polymer hydrogel capsules and cancer cell targeting. *ACS Nano.*2012;6:1463–72. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22260171)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=ACS+Nano&title=Macromolecule+functionalization+of+disulfide-bonded+polymer+hydrogel+capsules+and+cancer+cell+targeting&author=O+Shimoni&author=A+Postma&author=Y+Yan&author=AM+Scott&author=JK+Heath&volume=6&publication_year=2012&pages=1463-72&pmid=22260171&)]

16. Stumpel JE, Gil ER, Spoelstra AB, Bastiaansen CW, Broer DJ, Schenning AP. Stimuli-Responsive Materials Based on Interpenetrating Polymer Liquid Crystal Hydrogels. *Advanced Functional Materials.*2015;25:3314–20. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Functional+Materials&title=Stimuli-Responsive+Materials+Based+on+Interpenetrating+Polymer+Liquid+Crystal+Hydrogels&author=JE+Stumpel&author=ER+Gil&author=AB+Spoelstra&author=CW+Bastiaansen&author=DJ+Broer&volume=25&publication_year=2015&pages=3314-20&)]

17. Tanaka T. Collapse of gels and the critical endpoint. *Physical Review Letters.*1978;40:820. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Physical+Review+Letters&title=Collapse+of+gels+and+the+critical+endpoint&author=T+Tanaka&volume=40&publication_year=1978&pages=820&)]

18. Yatvin MB, Weinstein JN, Dennis WH, Blumenthal R. Design of liposomes for enhanced local release of drugs by hyperthermia. *Science.*1978;202:1290–3. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/364652)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Science&title=Design+of+liposomes+for+enhanced+local+release+of+drugs+by+hyperthermia&author=MB+Yatvin&author=JN+Weinstein&author=WH+Dennis&author=R+Blumenthal&volume=202&publication_year=1978&pages=1290-3&pmid=364652&)]

19. Mura S, Nicolas J, Couvreur P. Stimuli-responsive nanocarriers for drug delivery. *Nature Materials.*2013;12:991–1003. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/24150417)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nature+Materials&title=Stimuli-responsive+nanocarriers+for+drug+delivery&author=S+Mura&author=J+Nicolas&author=P+Couvreur&volume=12&publication_year=2013&pages=991-1003&pmid=24150417&)]

20. Kelley EG, Albert JN, Sullivan MO, Epps III TH. Stimuli-responsive copolymer solution and surface assemblies for biomedical applications. *Chemical Society Reviews.*2013;42:7057–71. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3703495/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23403471)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Chemical+Society+Reviews&title=Stimuli-responsive+copolymer+solution+and+surface+assemblies+for+biomedical+applications&author=EG+Kelley&author=JN+Albert&author=MO+Sullivan&author=TH+Epps+III&volume=42&publication_year=2013&pages=7057-71&pmid=23403471&)]

21. Liu J, Huang Y, Kumar A, Tan A, Jin S, Mozhi A. et al. pH-Sensitive nano-systems for drug delivery in cancer therapy. *Biotechnology Advances.*2014;32:693–710. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/24309541)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Biotechnology+Advances&title=pH-Sensitive+nano-systems+for+drug+delivery+in+cancer+therapy&author=J+Liu&author=Y+Huang&author=A+Kumar&author=A+Tan&author=S+Jin&volume=32&publication_year=2014&pages=693-710&pmid=24309541&)]

22. Ganesh VA, Baji A, Ramakrishna S. Smart functional polymers-a new route towards creating a sustainable environment. *RSC Advances.*2014;4:53352–64. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=RSC+Advances&title=Smart+functional+polymers-a+new+route+towards+creating+a+sustainable+environment&author=VA+Ganesh&author=A+Baji&author=S+Ramakrishna&volume=4&publication_year=2014&pages=53352-64&)]

23. Gao W, Chan JM, Farokhzad OC. pH-responsive nanoparticles for drug delivery. *Molecular Pharmaceutics.*2010;7:1913–20. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3379544/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/20836539)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Molecular+Pharmaceutics&title=pH-responsive+nanoparticles+for+drug+delivery&author=W+Gao&author=JM+Chan&author=OC+Farokhzad&volume=7&publication_year=2010&pages=1913-20&pmid=20836539&)]

24. Yu P, Yu H, Guo C, Cui Z, Chen X, Yin Q. et al. Reversal of doxorubicin resistance in breast cancer by mitochondria-targeted pH-responsive micelles. *Acta Biomaterialia.*2015;14:115–24. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/25498306)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Acta+Biomaterialia&title=Reversal+of+doxorubicin+resistance+in+breast+cancer+by+mitochondria-targeted+pH-responsive+micelles&author=P+Yu&author=H+Yu&author=C+Guo&author=Z+Cui&author=X+Chen&volume=14&publication_year=2015&pages=115-24&pmid=25498306&)]

25. Subudhi MB, Jain A, Jain A, Hurkat P, Shilpi S, Gulbake A. et al. Eudragit S100 coated citrus pectin nanoparticles for colon targeting of 5-Fluorouracil. *Materials.*2015;8:832–49. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5455456/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/28787974)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Materials&title=Eudragit+S100+coated+citrus+pectin+nanoparticles+for+colon+targeting+of+5-Fluorouracil&author=MB+Subudhi&author=A+Jain&author=A+Jain&author=P+Hurkat&author=S+Shilpi&volume=8&publication_year=2015&pages=832-49&)]

26. Stubbs M, McSheehy PM, Griffiths JR, Bashford CL. Causes and consequences of tumour acidity and implications for treatment. *Molecular Medicine Today.*2000;6:15–9. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/10637570)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Molecular+Medicine+Today&title=Causes+and+consequences+of+tumour+acidity+and+implications+for+treatment&author=M+Stubbs&author=PM+McSheehy&author=JR+Griffiths&author=CL+Bashford&volume=6&publication_year=2000&pages=15-9&pmid=10637570&)]

27. Neri D, Supuran CT. Interfering with pH regulation in tumours as a therapeutic strategy. *Nature Reviews Drug Discovery.*2011;10:767–77. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/21921921)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nature+Reviews+Drug+Discovery&title=Interfering+with+pH+regulation+in+tumours+as+a+therapeutic+strategy&author=D+Neri&author=CT+Supuran&volume=10&publication_year=2011&pages=767-77&pmid=21921921&)]

28. Lee ES, Oh KT, Kim D, Youn YS, Bae YH. Tumor pH-responsive flower-like micelles of poly (L-lactic acid)-b-poly (ethylene glycol)-b-poly (L-histidine) *Journal of Controlled Release.*2007;123:19–26. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2196406/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/17826863)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Journal+of+Controlled+Release&title=Tumor+pH-responsive+flower-like+micelles+of+poly+(L-lactic+acid)-b-poly+(ethylene+glycol)-b-poly+(L-histidine)&author=ES+Lee&author=KT+Oh&author=D+Kim&author=YS+Youn&author=YH+Bae&volume=123&publication_year=2007&pages=19-26&pmid=17826863&)]

29. Cheng R, Meng F, Deng C, Klok H-A, Zhong Z. Dual and multi-stimuli responsive polymeric nanoparticles for programmed site-specific drug delivery. *Biomaterials.*2013;34:3647–57. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23415642)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Biomaterials&title=Dual+and+multi-stimuli+responsive+polymeric+nanoparticles+for+programmed+site-specific+drug+delivery&author=R+Cheng&author=F+Meng&author=C+Deng&author=H-A+Klok&author=Z+Zhong&volume=34&publication_year=2013&pages=3647-57&pmid=23415642&)]

30. Pan Y-J, Chen Y-Y, Wang D-R, Wei C, Guo J, Lu D-R. et al. Redox/pH dual stimuli-responsive biodegradable nanohydrogels with varying responses to dithiothreitol and glutathione for controlled drug release. *Biomaterials.*2012;33:6570–9. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22704845)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Biomaterials&title=Redox/pH+dual+stimuli-responsive+biodegradable+nanohydrogels+with+varying+responses+to+dithiothreitol+and+glutathione+for+controlled+drug+release&author=Y-J+Pan&author=Y-Y+Chen&author=D-R+Wang&author=C+Wei&author=J+Guo&volume=33&publication_year=2012&pages=6570-9&pmid=22704845&)]

31. Chen W, Zhong P, Meng F, Cheng R, Deng C, Feijen J. et al. Redox and pH-responsive degradable micelles for dually activated intracellular anticancer drug release. *Journal of Controlled Release.*2013;169:171–9. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23306022)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Journal+of+Controlled+Release&title=Redox+and+pH-responsive+degradable+micelles+for+dually+activated+intracellular+anticancer+drug+release&author=W+Chen&author=P+Zhong&author=F+Meng&author=R+Cheng&author=C+Deng&volume=169&publication_year=2013&pages=171-9&pmid=23306022&)]

32. Huo M, Yuan J, Tao L, Wei Y. Redox-responsive polymers for drug delivery: from molecular design to applications. *Polymer Chemistry.*2014;5:1519–28. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Polymer+Chemistry&title=Redox-responsive+polymers+for+drug+delivery:+from+molecular+design+to+applications&author=M+Huo&author=J+Yuan&author=L+Tao&author=Y+Wei&volume=5&publication_year=2014&pages=1519-28&)]

33. Wang J, Sun X, Mao W, Sun W, Tang J, Sui M. et al. Tumor Redox Heterogeneity-Responsive Prodrug Nanocapsules for Cancer Chemotherapy. *Advanced Materials.*2013;25:3670–6. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23740675)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Materials&title=Tumor+Redox+Heterogeneity-Responsive+Prodrug+Nanocapsules+for+Cancer+Chemotherapy&author=J+Wang&author=X+Sun&author=W+Mao&author=W+Sun&author=J+Tang&volume=25&publication_year=2013&pages=3670-6&pmid=23740675&)]

34. Torchilin VP. Multifunctional, stimuli-sensitive nanoparticulate systems for drug delivery. *Nature Reviews Drug Discovery.*2014;13:813–27. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4489143/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/25287120)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nature+Reviews+Drug+Discovery&title=Multifunctional,+stimuli-sensitive+nanoparticulate+systems+for+drug+delivery&author=VP+Torchilin&volume=13&publication_year=2014&pages=813-27&pmid=25287120&)]

35. Wilson DS, Dalmasso G, Wang L, Sitaraman SV, Merlin D, Murthy N. Orally delivered thioketal nanoparticles loaded with TNF-α-siRNA target inflammation and inhibit gene expression in the intestines. *Nature Materials.*2010;9:923–8. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3142359/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/20935658)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nature+Materials&title=Orally+delivered+thioketal+nanoparticles+loaded+with+TNF-%CE%B1-siRNA+target+inflammation+and+inhibit+gene+expression+in+the+intestines&author=DS+Wilson&author=G+Dalmasso&author=L+Wang&author=SV+Sitaraman&author=D+Merlin&volume=9&publication_year=2010&pages=923-8&pmid=20935658&)]

36. Nguyen MM, Carlini AS, Chien MP, Sonnenberg S, Luo C, Braden RL. et al. Enzyme-Responsive Nanoparticles for Targeted Accumulation and Prolonged Retention in Heart Tissue after Myocardial Infarction. *Advanced Materials.*2015;27:5547–52. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4699559/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/26305446)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Materials&title=Enzyme-Responsive+Nanoparticles+for+Targeted+Accumulation+and+Prolonged+Retention+in+Heart+Tissue+after+Myocardial+Infarction&author=MM+Nguyen&author=AS+Carlini&author=MP+Chien&author=S+Sonnenberg&author=C+Luo&volume=27&publication_year=2015&pages=5547-52&pmid=26305446&)]

37. Callmann CE, Barback CV, Thompson MP, Hall DJ, Mattrey RF, Gianneschi NC. Therapeutic Enzyme-Responsive Nanoparticles for Targeted Delivery and Accumulation in Tumors. *Advanced Materials.*2015;27:4611–5. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4699560/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/26178920)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Materials&title=Therapeutic+Enzyme-Responsive+Nanoparticles+for+Targeted+Delivery+and+Accumulation+in+Tumors&author=CE+Callmann&author=CV+Barback&author=MP+Thompson&author=DJ+Hall&author=RF+Mattrey&volume=27&publication_year=2015&pages=4611-5&pmid=26178920&)]

38. De La Rica R, Aili D, Stevens MM. Enzyme-responsive nanoparticles for drug release and diagnostics. *Advanced Drug Delivery Reviews.*2012;64:967–78. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22266127)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Drug+Delivery+Reviews&title=Enzyme-responsive+nanoparticles+for+drug+release+and+diagnostics&author=R+De+La+Rica&author=D+Aili&author=MM+Stevens&volume=64&publication_year=2012&pages=967-78&pmid=22266127&)]

39. [39] Lock LL, Tang Z, Keith D, Reyes C, Cui H. Enzyme-Specific Doxorubicin Drug Beacon as Drug-Resistant Theranostic Molecular Probes. *ACS Macro Letters.*2015;4:552–5. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=ACS+Macro+Letters&title=Enzyme-Specific+Doxorubicin+Drug+Beacon+as+Drug-Resistant+Theranostic+Molecular+Probes&author=LL+%5b39%5d+Lock&author=Z+Tang&author=D+Keith&author=C+Reyes&author=H+Cui&volume=4&publication_year=2015&pages=552-5&)]

40. Shi Y, van den Dungen ET, Klumperman B, van Nostrum CF, Hennink WE. Reversible Addition-Fragmentation Chain Transfer Synthesis of a Micelle-Forming, Structure Reversible Thermosensitive Diblock Copolymer Based on the N-(2-Hydroxy propyl) Methacrylamide Backbone. *ACS Macro Letters.*2013;2:403–8. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=ACS+Macro+Letters&title=Reversible+Addition-Fragmentation+Chain+Transfer+Synthesis+of+a+Micelle-Forming,+Structure+Reversible+Thermosensitive+Diblock+Copolymer+Based+on+the+N-(2-Hydroxy+propyl)+Methacrylamide+Backbone&author=Y+Shi&author=ET+van+den+Dungen&author=B+Klumperman&author=CF+van+Nostrum&author=WE+Hennink&volume=2&publication_year=2013&pages=403-8&)]

41. Shi Y, van Steenbergen MJ, Teunissen EA, Novo Ls, Gradmann S, Baldus M. et al. Π-Π stacking increases the stability and loading capacity of thermosensitive polymeric micelles for chemotherapeutic drugs. *Biomacromolecules.*2013;14:1826–37. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23607866)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Biomacromolecules&title=%CE%A0-%CE%A0+stacking+increases+the+stability+and+loading+capacity+of+thermosensitive+polymeric+micelles+for+chemotherapeutic+drugs&author=Y+Shi&author=MJ+van+Steenbergen&author=EA+Teunissen&author=Ls+Novo&author=S+Gradmann&volume=14&publication_year=2013&pages=1826-37&pmid=23607866&)]

42. Shi Y, Cardoso RM, Van Nostrum CF, Hennink WE. Anthracene functionalized thermosensitive and UV-crosslinkable polymeric micelles. *Polymer Chemistry.*2015;6:2048–53. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Polymer+Chemistry&title=Anthracene+functionalized+thermosensitive+and+UV-crosslinkable+polymeric+micelles&author=Y+Shi&author=RM+Cardoso&author=CF+Van+Nostrum&author=WE+Hennink&volume=6&publication_year=2015&pages=2048-53&)]

43. Danhier F, Feron O, Préat V. To exploit the tumor microenvironment: passive and active tumor targeting of nanocarriers for anti-cancer drug delivery. *Journal of Controlled Release.*2010;148:135–46. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/20797419)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Journal+of+Controlled+Release&title=To+exploit+the+tumor+microenvironment:+passive+and+active+tumor+targeting+of+nanocarriers+for+anti-cancer+drug+delivery&author=F+Danhier&author=O+Feron&author=V+Pr%C3%A9at&volume=148&publication_year=2010&pages=135-46&pmid=20797419&)]

44. Adelsberger J, Kulkarni A, Jain A, Wang W, Bivigou-Koumba AM, Busch P. et al. Thermoresponsive PS-b-PNIPAM-b-PS micelles: aggregation behavior, segmental dynamics, and thermal response. *Macromolecules.*2010;43:2490–501. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Macromolecules&title=Thermoresponsive+PS-b-PNIPAM-b-PS+micelles:+aggregation+behavior,+segmental+dynamics,+and+thermal+response&author=J+Adelsberger&author=A+Kulkarni&author=A+Jain&author=W+Wang&author=AM+Bivigou-Koumba&volume=43&publication_year=2010&pages=2490-501&)]

45. Zhao Y, Fan X, Liu D, Wang Z. PEGylated thermo-sensitive poly (amidoamine) dendritic drug delivery systems. *International Journal of Pharmaceutics.*2011;409:229–36. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/21316434)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=International+Journal+of+Pharmaceutics&title=PEGylated+thermo-sensitive+poly+(amidoamine)+dendritic+drug+delivery+systems&author=Y+Zhao&author=X+Fan&author=D+Liu&author=Z+Wang&volume=409&publication_year=2011&pages=229-36&pmid=21316434&)]

46. Lal S, Clare SE, Halas NJ. Nanoshell-enabled photothermal cancer therapy: impending clinical impact. *Accounts of Chemical Research.*2008;41:1842–51. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/19053240)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Accounts+of+Chemical+Research&title=Nanoshell-enabled+photothermal+cancer+therapy:+impending+clinical+impact&author=S+Lal&author=SE+Clare&author=NJ+Halas&volume=41&publication_year=2008&pages=1842-51&pmid=19053240&)]

47. Sun J, Zhang Y, Chen Z, Zhou J, Gu N. Fibrous Aggregation of Magnetite Nanoparticles Induced by a Time-Varied Magnetic Field. *Angewandte Chemie International Edition.*2007;46:4767–70. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/17506053)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Angewandte+Chemie+International+Edition&title=Fibrous+Aggregation+of+Magnetite+Nanoparticles+Induced+by+a+Time-Varied+Magnetic+Field&author=J+Sun&author=Y+Zhang&author=Z+Chen&author=J+Zhou&author=N+Gu&volume=46&publication_year=2007&pages=4767-70&)]

48. Liu J, Zhang Y, Wang C, Xu R, Chen Z, Gu N. Magnetically sensitive alginate-templated polyelectrolyte multilayer microcapsules for controlled release of doxorubicin. *The Journal of Physical Chemistry C.*2010;114:7673–9. [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=The+Journal+of+Physical+Chemistry+C&title=Magnetically+sensitive+alginate-templated+polyelectrolyte+multilayer+microcapsules+for+controlled+release+of+doxorubicin&author=J+Liu&author=Y+Zhang&author=C+Wang&author=R+Xu&author=Z+Chen&volume=114&publication_year=2010&pages=7673-9&)]

49. Chen Z, Yin J-J, Zhou Y-T, Zhang Y, Song L, Song M. et al. Dual enzyme-like activities of iron oxide nanoparticles and their implication for diminishing cytotoxicity. *Acs Nano.*2012;6:4001–12. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22533614)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Acs+Nano&title=Dual+enzyme-like+activities+of+iron+oxide+nanoparticles+and+their+implication+for+diminishing+cytotoxicity&author=Z+Chen&author=J-J+Yin&author=Y-T+Zhou&author=Y+Zhang&author=L+Song&volume=6&publication_year=2012&pages=4001-12&pmid=22533614&)]

50. Fang K, Song L, Gu Z, Yang F, Zhang Y, Gu N. Magnetic field activated drug release system based on magnetic PLGA microspheres for chemo-thermal therapy. *Colloids and Surfaces B: Biointerfaces.*2015;136:712–20. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/26513754)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Colloids+and+Surfaces+B:+Biointerfaces&title=Magnetic+field+activated+drug+release+system+based+on+magnetic+PLGA+microspheres+for+chemo-thermal+therapy&author=K+Fang&author=L+Song&author=Z+Gu&author=F+Yang&author=Y+Zhang&volume=136&publication_year=2015&pages=712-20&pmid=26513754&)]

51. Yang F, Zhang X, Song L, Cui H, Myers JN, Bai T. et al. Controlled Drug Release and Hydrolysis Mechanism of Polymer-Magnetic Nanoparticle Composite. *ACS Applied Materials & Interfaces.*2015;7:9410–9. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/25881356)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=ACS+Applied+Materials+&+Interfaces&title=Controlled+Drug+Release+and+Hydrolysis+Mechanism+of+Polymer-Magnetic+Nanoparticle+Composite&author=F+Yang&author=X+Zhang&author=L+Song&author=H+Cui&author=JN+Myers&volume=7&publication_year=2015&pages=9410-9&pmid=25881356&)]

52. Hu K, Sun J, Guo Z, Wang P, Chen Q, Ma M. et al. A novel magnetic hydrogel with aligned magnetic colloidal assemblies showing controllable enhancement of magnetothermal effect in the presence of alternating magnetic field. *Advanced Materials.*2015;27:2507–14. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/25753892)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Advanced+Materials&title=A+novel+magnetic+hydrogel+with+aligned+magnetic+colloidal+assemblies+showing+controllable+enhancement+of+magnetothermal+effect+in+the+presence+of+alternating+magnetic+field&author=K+Hu&author=J+Sun&author=Z+Guo&author=P+Wang&author=Q+Chen&volume=27&publication_year=2015&pages=2507-14&pmid=25753892&)]

53. Wang F, Kim D-K, Yoshitake T, Johansson S, Bjelke B, Muhammed M. et al. Diffusion and clearance of superparamagnetic iron oxide nanoparticles infused into the rat striatum studied by MRI and histochemical techniques. *Nanotechnology.*2010;22:015103. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/21135466)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nanotechnology&title=Diffusion+and+clearance+of+superparamagnetic+iron+oxide+nanoparticles+infused+into+the+rat+striatum+studied+by+MRI+and+histochemical+techniques&author=F+Wang&author=D-K+Kim&author=T+Yoshitake&author=S+Johansson&author=B+Bjelke&volume=22&publication_year=2010&pages=015103&pmid=21135466&)]

54. Yue-Jian C, Juan T, Fei X, Jia-Bi Z, Ning G, Yi-Hua Z. et al. Synthesis, self-assembly, and characterization of PEG-coated iron oxide nanoparticles as potential MRI contrast agent. *Drug Development and Industrial Pharmacy.*2010;36:1235–44. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/20818962)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Drug+Development+and+Industrial+Pharmacy&title=Synthesis,+self-assembly,+and+characterization+of+PEG-coated+iron+oxide+nanoparticles+as+potential+MRI+contrast+agent&author=C+Yue-Jian&author=T+Juan&author=X+Fei&author=Z+Jia-Bi&author=G+Ning&volume=36&publication_year=2010&pages=1235-44&pmid=20818962&)]

55. Xie J, Zhang Y, Yan C, Song L, Wen S, Zang F. et al. High-performance PEGylated Mn-Zn ferrite nanocrystals as a passive-targeted agent for magnetically induced cancer theranostics. *Biomaterials.*2014;35:9126–36. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/25106772)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Biomaterials&title=High-performance+PEGylated+Mn-Zn+ferrite+nanocrystals+as+a+passive-targeted+agent+for+magnetically+induced+cancer+theranostics&author=J+Xie&author=Y+Zhang&author=C+Yan&author=L+Song&author=S+Wen&volume=35&publication_year=2014&pages=9126-36&pmid=25106772&)]

56. Xiong F, Chen Y, Chen J, Yang B, Zhang Y, Gao H. et al. Rubik-like magnetic nanoassemblies as an efficient drug multifunctional carrier for cancer theranostics. *Journal of Controlled Release.*2013;172:993–1001. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/24096016)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Journal+of+Controlled+Release&title=Rubik-like+magnetic+nanoassemblies+as+an+efficient+drug+multifunctional+carrier+for+cancer+theranostics&author=F+Xiong&author=Y+Chen&author=J+Chen&author=B+Yang&author=Y+Zhang&volume=172&publication_year=2013&pages=993-1001&pmid=24096016&)]

57. Song L, Zang F, Song M, Chen G, Zhang Y. Effective PEGylation of Fe3O4 nanomicelles for in vivo MR imaging. *Journal of Nanoscience and Nanotechnology.*2015;15:4111–8. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/26369019)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Journal+of+Nanoscience+and+Nanotechnology&title=Effective+PEGylation+of+Fe3O4+nanomicelles+for+in+vivo+MR+imaging&author=L+Song&author=F+Zang&author=M+Song&author=G+Chen&author=Y+Zhang&volume=15&publication_year=2015&pages=4111-8&pmid=26369019&)]

58. Liu D, Wu W, Chen X, Wen S, Zhang X, Ding Q. et al. Conjugation of paclitaxel to iron oxide nanoparticles for tumor imaging and therapy. *Nanoscale.*2012;4:2306–10. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/22362270)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Nanoscale&title=Conjugation+of+paclitaxel+to+iron+oxide+nanoparticles+for+tumor+imaging+and+therapy&author=D+Liu&author=W+Wu&author=X+Chen&author=S+Wen&author=X+Zhang&volume=4&publication_year=2012&pages=2306-10&pmid=22362270&)]

59. Yang H-W, Hua M-Y, Liu H-L, Huang C-Y, Tsai R-Y, Lu Y-J. et al. Self-protecting core-shell magnetic nanoparticles for targeted, traceable, long half-life delivery of BCNU to gliomas. *Biomaterials.*2011;32:6523–32. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/21645920)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Biomaterials&title=Self-protecting+core-shell+magnetic+nanoparticles+for+targeted,+traceable,+long+half-life+delivery+of+BCNU+to+gliomas&author=H-W+Yang&author=M-Y+Hua&author=H-L+Liu&author=C-Y+Huang&author=R-Y+Tsai&volume=32&publication_year=2011&pages=6523-32&pmid=21645920&)]

60. Hayashi K, Nakamura M, Sakamoto W, Yogo T, Miki H, Ozaki S. et al. Superparamagnetic nanoparticle clusters for cancer theranostics combining magnetic resonance imaging and hyperthermia treatment. *Theranostics.*2013;3:366–76. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3677408/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23781284)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Theranostics&title=Superparamagnetic+nanoparticle+clusters+for+cancer+theranostics+combining+magnetic+resonance+imaging+and+hyperthermia+treatment&author=K+Hayashi&author=M+Nakamura&author=W+Sakamoto&author=T+Yogo&author=H+Miki&volume=3&publication_year=2013&pages=366-76&pmid=23781284&)]

61. Paris JL, Cabañas MV, Manzano M, Vallet-Regí M. Polymer-Grafted Mesoporous Silica Nanoparticles as Ultrasound-Responsive Drug Carriers. *ACS Nano.*2015;9:11023–33. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/26456489)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=ACS+Nano&title=Polymer-Grafted+Mesoporous+Silica+Nanoparticles+as+Ultrasound-Responsive+Drug+Carriers&author=JL+Paris&author=MV+Caba%C3%B1as&author=M+Manzano&author=M+Vallet-Reg%C3%AD&volume=9&publication_year=2015&pages=11023-33&pmid=26456489&)]

62. Guo Q, Zhang T, An J, Wu Z, Zhao Y, Dai X. et al. Block versus Random Amphiphilic Glycopolymer Nanopaticles as Glucose-Responsive Vehicles. *Biomacromolecules.*2015;16:3345–56. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/26397308)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Biomacromolecules&title=Block+versus+Random+Amphiphilic+Glycopolymer+Nanopaticles+as+Glucose-Responsive+Vehicles&author=Q+Guo&author=T+Zhang&author=J+An&author=Z+Wu&author=Y+Zhao&volume=16&publication_year=2015&pages=3345-56&pmid=26397308&)]

63. Wu Q, Wang L, Yu H, Wang J, Chen Z. Organization of glucose-responsive systems and their properties. *Chemical Reviews.*2011;111:7855–75. [[PubMed](https://pubmed.ncbi.nlm.nih.gov/21902252)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=Chemical+Reviews&title=Organization+of+glucose-responsive+systems+and+their+properties&author=Q+Wu&author=L+Wang&author=H+Yu&author=J+Wang&author=Z+Chen&volume=111&publication_year=2011&pages=7855-75&pmid=21902252&)]

64. Gu Z, Aimetti AA, Wang Q, Dang TT, Zhang Y, Veiseh O. et al. Injectable nano-network for glucose-mediated insulin delivery. *ACS Nano.*2013;7:4194–201. [[PMC free article](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4107450/)] [[PubMed](https://pubmed.ncbi.nlm.nih.gov/23638642)] [[Google Scholar](https://scholar.google.com/scholar_lookup?journal=ACS+Nano&title=Injectable+nano-network+for+glucose-mediated+insulin+delivery&author=Z+Gu&author=AA+Aimetti&author=Q+Wang&author=TT+Dang&author=Y+Zhang&volume=7&publication_year=2013&pages=4194-201&pmid=23638642&)]