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Electrochemical fabrication of mesoporous metalalloy films

Javeria Bashir, (10)†ab Md. Belal Chowdhury, (10)†c Rahanuma Raihanu Kathak,†c Shuvashis Dey, (10) Atiya Tahira Tasnim, Mohammed A. Amin, Yusuf Valentino Kaneti, Db Mostafa Kamal Masud D*bce and Md. Shahriar A Hossain ** **ab

Nanoarchitectured mesoporous metal alloy films offer immense potential by integrating the intrinsic catalytic capabilities of their constituent metals to create suitable surface morphologies, different signal transductions and catalytic phenomena. These unique properties and functionalities have led to the development of several strategies for fabricating mesoporous alloy films, including template-based (e.g., lyotropic liquid crystalline (LLC) and di-block polymer-based templating) and template-free approaches (e.g., post-synthetic dealloying). More recently, the utilization of electrochemical methods for metallic mesoporous alloy films has ushered in a new way of fabrication and attracted much attention due to their simplicity, environmental friendliness, and cost-effectiveness. In this review article, we comprehensively reviewed the key electrochemical approaches for fabricating various types of metallic mesoporous alloy films (a wide range of geometries, pore sizes, form and tunneling structures, and metal compositions). With an emphasis on the fabrication of mesoporous bimetallic and trimetallic alloy films, we discuss their specific application for developing advanced technologies spanning from hydrogen production to fuel cell and sensor development. Furthermore, we also discuss the bottlenecks in the fabrication of mesoporous alloy films and provide a future perspective on the techniques. We envisage that this literature will guide the researcher towards research exploring engineered alloying of earth-abundant elements and translating them for industrial-scale electrocatalysis and nanomedicine application.

1. Introduction

Mesoporous nanomaterials including nanoparticles, films and building blocks have gained immense interest in recent years owing to their porous architecture that provides an exceptionally large surface area, mesochannels, and enhanced mass and ion transfer inside a nano-level material. These unique and fascinating features make them highly favorable and suitable for widespread applications including adsorption, isolation and separation of large molecules including chemicals and

biomolecules,2 catalysis,3,4 targeted drug and nucleic acid delivery,3,5-9 sensor fabrication,10 advanced functional devices^{11,12} and highly efficient fuel cells.¹³ As a result, extensive research has been conducted to develop highly efficient mesoporous materials based on the target application, leading to the exploration of a wide range of mesoporous nanostructures of various chemical nature (e.g., carbon, metals, silica, aluminosilicates, metal oxides, metal carbides, metal sulphides, metal nitrides, polymers, etc.) as well as various frameworks (e.g., nanomaterials, nanocomposites and films). 14 Among these materials, mesoporous metal architectures, particularly mesoporous metal films, are fascinating due to their intriguing physiochemical properties, such as ultra-large surface area, numerous exposed catalytic active sites, a widely open and interconnected porous structure that provides fast mass transport, high electrical conductivity, high mechanical stability and the ability to functionalize the pore surface in a variety of ways. 15-18 Simultaneously, the ever-increasing demand for new materials with desirable properties and functionalities has fueled research into the synthesis of metallic alloys and fabrication of relevant devices. 19-24

Mesoporous metal alloy architectures combine the intrinsic catalytic characteristics of their constituent metals with the

^a School of Mechanical and Mining Engineering, Faculty of Engineering, Architecture and Information Technology (EAIT), The University of Queensland, Brisbane, QLD 40721, Australia. E-mail: md.hossainn@uq.edu.au

^b Australian Institute for Bioengineering and Nanotechnology (AIBN), The University of Queensland, Brisbane, QLD 4072, Australia. E-mail: m.masud@uq.edu.au

^c Department of Biochemistry and Molecular Biology, School of Life Sciences, Shahjalal University of Science & Technology, Sylhet 3114, Bangladesh

^d Department of Chemistry, College of Science, Taif University, P.O. Box 11099, Taif, 21944, Saudi Arabia

^e JST-ERATO Yamauchi Materials Space-Tectonics Project and International Center for Materials Nanoarchitectonics (WPI-MANA), National Institute for Materials Science, Tsukuba, Ibaraki 305-0044, Japan

[†] These authors contributed equally.

unique properties of mesoporous structures to provide distinctive signal transduction and catalytic properties. Mesoporous alloy films made up of multiple metals with varied reactivities and standard redox potentials (E^0) can provide several advantages for electrocatalysis. 21,23 Compared to the traditional nonporous noble metal based electrocatalysts, mesoporous metal alloy nanoarchitectures with a combination of noble metals can enhance electrocatalytic performance significantly because they produce a more favorable electrical structure and reduce the electrode fabrication cost. For instance, mesoporous Au-Cu films exhibit excellent performance in developing nonenzymatic glucose sensors with high sensitivity and selectivity, and reduce the cost involved in device fabrication due to the synergism between electronic response and the catalytic benefits from both components compared to that of monometallic Au based devices. 23 Such superior electrocatalytic performances have also been reported for other mesoporous films, e.g., Pt-Cu,²⁵ Au-Ni,²⁶ Rh-Ni,²¹ etc. Moreover, mesoporous noble

metal-metal mixtures (e.g., Au-Pt, Pt-Pd, 24,25 Pt-Ru, 27 Au-Ag, Au-Ni,26 etc.) have resulted in the development of highperformance heterogeneous electrocatalysts with better durability. For instance, the catalytic activity and electrochemical stability of mesoporous Pt-Au alloys or Pt-Ru alloys towards methanol oxidation (methanol fuel cell) and/or glucose oxidation (glucose detection) are substantially greater than that of mesoporous Pt films, which can be attributed to the welldeveloped mesoporous structure with a large surface area and the synergistic characteristics of Pt and Au. 24,27

To date, several strategies for fabricating mesoporous alloy films have been reported, which can be classified as the hardtemplating approach (i.e., nanocasting),29 the soft-templating approach (e.g., lyotropic liquid crystalline (LLC) and di-block polymer based templating)^{30,31} and the template-free approach (e.g., post-synthetic dealloying).³² Hard templates made of inorganic materials such as mesoporous silica are commonly used to guide and control the building of mesoporous metals.



Iaveria Bashir

Iaveria Bashir is a PhD student at the Australian Institute for Bioengineering and Nanotechnology (AIBN), The University of Oueensland Australia. (UQ), She received her master's degree in physics (2020) from the University of Lahore, Pakistan. Her research focuses on the synthesis and manipulation of magneto-plasmonic biocompatible mesoporous nanostructures for SERS based biosensing of clinically relevant biomarkers.



Md. Belal Chowdhury

Md. Belal Chowdhury graduated from the Department of Biochemistry and Molecular Biology from Shahjalal University of Science and Technology, Sylhet, Bangladesh. His research interest centers on the development of nanoarchitectured nanoparticles for early detection of diseasespecific biomarkers with high sensitivity, specificity, and reliability along with faster detection time compared to the conventional diagnostic solutions.

He has received several honors and distinctions for his excellent track record in academic study and published 4 peer-reviewed original research articles.



Rahanuma Raihanu Kathak

Rahanuma Raihanu Kathak is a lecturer of Biochemistry and Molecular Biology (BMB) at Primeasia University, Bangladesh. She has received her Master of Science and Bachelor of Science degrees in BMB from Shahjalal University of Science and Tech-(SUST), *Sylhet-3114*, nology Bangladesh, with an academic record of 3.99 and 3.85 (out of 4.00) respectively. Her research interest lies in the field of medical diagnostics, especially

in the emerging application of nano-architectonics for early disease detection.



Shuvashis Dev

Dr Shuvashis Dey is a Postdoctoral Research Fellow at the Australian Institute for Bioengineering and Nanotechnology (AIBN), The University of Queensland (UQ). Prior to joining his current position, he received his PhD degree in biomedical engineering from UQ and developed microfluidic biosensors for early diagnosis and personalized treatment for better patient management. His research interest includes design and fabrication

of micro/nanofluidic platforms and utilization of plasmonic nanomaterials for biosensing purpose.

The generated replicas reverse the template's morphology; however, the dissolution process is typically slow and requires repeated exposure to dangerous chemicals. Although hardtemplates provide a consistent and intuitive method for producing mesoporous metals, the complexity of the technique, lack of morphology control, and use of harsh chemicals have confined their use in wider applications.^{9,33} Soft-templating methods are promising alternatives that use organic surfactants (e.g., block polymers, lyotropic liquid crystals, etc.) as molecular templates to direct the formation of mesopores in metallic structures.31 Though soft-templating methods offer several distinctive benefits (e.g., cost-effectiveness, a wider porous structure with designed pore size and morphology, and customization), the high viscosity of the formed micellar system limits their applications.³⁴ Mesoporous metal nano-

structures have also been synthesized using a dealloying technique that selectively removes the most chemically active element.³⁵ The chemical treatment preferentially dissolves a non-targeted element, leaving a porous metal or alloy surface behind. This technique is commonly seen in technologically significant alloy systems, such as stainless steel, brass, and different alloys especially Cu-Al alloys.³⁶ However, the mechanical and physical properties of a porous overlayer differ greatly from those of the bulk alloy on which it sits, resulting in stress corrosion cracking, brittle crack propagation as well as other unwanted material failure. 37 Even though these approaches are widely used for fabricating alloyed mesoporous metal architectures, these methods are complex and involve multistep processes, limiting their potential applicability, flexibility, and reproducibility. To address these issues, research has



Atiya Tahira Tasnim

Atiya Tahira Tasnim is a graduate student at the Department of Biochemistry and Molecular Biology under the School of Life Sciences at Shahjalal Univerity of Science and Technology, Bangladesh. Her research interest focuses on the scope of biomolecule-based cancer nanomedicine, developing therapeutics in silico with a combination of wet-lab techniques, and exploring cutting-edge biotechnological methods ranging from single-cell genomics to nanotechnology.



Yusuf Valentino Kaneti

Valentino Yusuf Kaneti received his PhD from the University of New South Wales in 2014. He has conducted postdoctoral research at Monash University (Australia), Graduate School at Shenzhen, Tsinghua University (China), and National Institute for Materials Science (Japan). Currently, he is working as an Advance Queensland Industry Research Fellow at the Australian Institute for Bioengineering and Nanotechnology, The University

of Queensland. His research focuses on the rational design and fabrication of functional nanoporous materials and hybrid materials for energy and environmental applications.



Mostafa Kamal Masud

Dr Mostafa Kamal Masud is a Postdoctoral Research Fellow at Australian Institute for Bioengineering and Nanotechnology (AIBN), TheUniversity Queensland. Prior to joining AIBN, he has worked as a JSPS postdoctoral fellow at National Institute for Materials Science, Japan. He obtained his PhD The University of Queensland, Australia, in 2020. His research focuses on the design and translation of bio-favourable

engineered mesoporous nanostructures for the development of nanobiosensor to address critical issues in medical diagnosis.



Md. Shahriar A Hossain

Dr Shahriar Hossain is an Associate Professor at the School of Mechanical and Mining Engineering at The University of Queensland, Australia. He obtained his PhD degree in Materials Science and Engineering from the Institute for Superconducting and Electronic Materials (ISEM) at University of Wollongong (UoW), Australia, in 2008. Dr Hossain was then employed as a Postdoctoral Research Fellow at the

University of Geneva (Switzerland). He was awarded the prestigious DECRA Fellowship from ARC in 2013. His research expertise is on the fabrication and characterization of superconducting materials and magnetically triggered nanoparticles for medical, power, and energy applications.

concentrated on the development of new strategies for the synthesis of more effective mesoporous materials in terms of fabrication time, flexibility, electrical properties, catalytic performance, and expense. In this regard, electrochemistry has opened an excellent and important window for the mesoporous world in terms of both electrochemical fabrication and characterization of mesoporous metal-based electrode structures. 17,38,39 Compared to other methods, electrochemical synthesis of mesoporous alloy films is considered as a competitive and versatile synthetic method, in terms of simplicity, environmental friendliness and cost-effectiveness. 14,22-26,40 The basic working principle of electrochemical techniques involves the application of an external potential on an appropriate electrode (as a working electrode) to convert the soluble precursor chemicals present in the electrolyte solution into solid mesoporous films. Metallic alloys have frequently been fabricated on conductive (as an electrode) surfaces using only one operational step, that is the simultaneous reduction of the precursor ions contained in the electrolyte by adjusting the potential/current density.41 A distinguishing feature of electrodeposition is that several reactions can occur simultaneously at the same working electrode that can be stimulated based on the external potential used.¹⁶ Because the applied potential is the driving force for depositing metal precursors dissolved in the electrolyte to form alloy films, the thickness of electrodeposited films can be controlled by varying the deposition time and metal precursor composition present in the electrolyte. Furthermore, because the films are deposited on conductive substrates, the as-prepared film can be immediately used for application purposes. 20,21,24,42,43 Even though a plethora of research has been done in this field, no reviews have yet been published on the fabrication of mesoporous metal alloys utilizing both direct and indirect strategies, as well as on their applications. To address the research gap and to lead future science in mesoporous alloy films and their application in catalysis, energy, chemical sensor and biosensor applications, reports are sought in which both the fabrication and film-based targeted application are

In this review we have comprehensively focused on the various electrochemical approaches for synthesizing different types (a wide range of geometries, pore sizes, shape and tunneling structures, and metal compositions) of metallic mesoporous alloy films. With an emphasis on the fabrication of mesoporous bimetallic and trimetallic alloy films, we have discussed their specific application for developing advanced technologies spanning from hydrogen production to fuel cell and sensor development. Following a thorough discussion in each section, this review also chronicles the challenges and needs implicated in the fabrication of mesoporous alloy films and provides a future perspective on the strategies presented.

2. Strategies for fabricating mesoporous metal alloy films

comprehensively discussed.

With the growing interest in mesoporous metal alloys with controllable porosity and pore morphology (size, shape, distribution

and tunneling) as well as unique chemical and physical properties, several strategies have been developed to obtain the porous alloy film of interest. The cutting-edge nanotechnology is capable of developing porous structures with pore sizes ranging from 2 nm to 100 nm and spacing widths ranging from 50 to 100 nm with desired geometry.44 The most used strategies adapted for fabricating such alloys include templatebased strategies (soft and hard template), potential-controlled surfactant assembly, colloidal assembly, interfacial surfactant templating, deposition from lyotropic liquid crystalline phases of surfactants, sol-gel assembly, electro-assisted self-assembly method, etc. Each strategy is specific and provides films with defined shape, quality, elemental composition, architecture, and pore definition. The pore size, shape, and morphology are also affected by the substrate (e.g., silicon wafer, glass, stainless steel, etc.) and template used to develop porous structures. For instance, synthesis of mesoporous titania films on freshly ground steel produced extensively fractured films with distorted porosity, whereas films produced on pre-calcined steel demonstrated good integrity, high substrate coverage, and a narrow pore size distribution with ordered and interconnected pores. 45 Though the above mentioned strategies have their own advantages and superiority, each methodology also suffers from drawbacks. 27,46,47 Among all reported strategies, the template based strategies (e.g. endo-template or soft-templating approach, exo-template or hard templating approach, etc.)48-51 are popular where an engineered template is used to direct the geometry of pores in the alloy films. Herein, we have divided the above-mentioned strategies into two broad categories: (i) direct electrodeposition of metals by electroreduction of metal cations, and (ii) indirect precipitation of metal hydroxides through electrogenerated hydroxide ions.

2.1. Direct electrodeposition of metals by electroreduction of metal cations

This is one of the most common deposition processes that involves the electrochemical reduction of metal precursors to their solid metal states, where electrons are directly transferred by the working electrode. 39,50 In the direct electrodeposition, the metal ions interact with the polar part of polymeric micelles. The micelles' geometry depends on the nature of both free ions and the polymer. These ions as well as the ion-capped polymeric micelles become positively charged even though initially they are non-ionic. If some paramagnetic ions like Ni²⁺ are present in the micellar system, they have no significant effect on the process in the absence of a magnetic field. However, in the presence of an electric field, the capped micelles move to the negative electrode where the ion precursors are reduced. Therefore, the applied electric field is a critical factor for obtaining the desired porous structure. The reduction potential is also a determining factor of the porous structure as during the deposition process, a higher reduction rate is achieved through a large negative potential. If two precursors with different potential are present in the system, the precursor having higher redox potential (value vs. the standard electrode) will be favored for reduction. For example,

at a deposition potential of -0.7 V (vs. Ag/AgCl), the reduction of the Au precursor in micelles containing both Au and Ni ions is higher due to the large difference between the standard redox potentials of Au and Ni (i.e., the standard redox potential of [AuCl₄]⁻/Au is 1.00 V vs. standard calomel electrode (SCE), while the standard redox potential of Ni^{2+}/Ni is -0.25 V vs. SCE). Within a certain range of applied potential the metal alloy film can be fabricated but no porosity can be observed due to too fast or too slow reduction of ions. Therefore, to get highly uniform and porous metal alloy films, the voltage always needs to be optimized.²⁸ Overall, the electrical potential applied to electrodes regulates the rate of electron generation and, as a result, nucleation and crystal growth. Temperature also plays a big role as higher temperature leads to large ligament of the mesoporous structure having large size nanocrystals. These variables are critical in determining the morphology of the nanoporous film. These electrodeposition techniques have grown in popularity over the last two decades as a means of developing efficient and reliable processes for creating meso-

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porous metals.16

Soft template approach. In current nanoscience, the soft templates are used as pore directors (e.g., block copolymers, surfactants, supramolecular assembly and aggregates) in the synthesis of silica, organo-silica, aluminosilicates, metals, polymers and phosphates.⁵² The inorganic precursors (metal salts of interest) react in a non-homogeneous medium (a solution containing one of the pore directing agents) in such a way that the shape of the final solid material follows the dynamic structure of the soft template that functions as a nano or microreactor. Soft systems are typically composed of two components, the template and the metal precursors that are dissolved in an aqueous component.⁵³ The major benefits of the soft template approach include (i) high control over the mesostructure and the size of pores by directing the adjustable structure of the prepared soft systems; (ii) simplicity and cost effectiveness due to fewer synthetic steps in comparison to hard templating approaches; (iii) ability to obtain wide and varied pore morphologies; (iv) applicability to produce mesoporous materials on a large scale; (v) synthesis of high quality mesoporous materials; and (vi) avoidance of the washing procedure and the use of toxic chemicals. However, there are some disadvantages with this approach, which are (i) difficulty in removing the surfactants and the adsorbed organic species; (ii) use of surfactants; (iii) high reactivity and sensitivity, which can affect the mesostructure; and (iv) low crystallinity of synthesized materials. 52-57 However, soft templating remains challenging due to the crystal growth which is difficult to

Hard template approach. The hard template-based strategy has been considered as one of the most powerful strategies for the fabrication of mesoporous materials. During the synthesis, the metal ion deposition takes place in the holes of the porous solid material which acts as a hard template, to give the final solid material.⁵⁸ The hard template approach has been widely used to synthesize a variety of porous materials in the last few decades, most notably those based on carbon, metals, metal

sulphides, metal oxides, metal carbides, metal nitrides, nonoxide ceramics and zeolites.⁵⁹ This strategy is also known as nanocasting as the process requires filling a hard template to achieve a negative replica of the rigid template to obtain a defined porous structure.33 This approach only uses rigid templates. The notable benefits of this approach are (i) high quality mesoporous materials in terms of crystallinity and porosity and (ii) the relatively high independence and insensibility of mesoporous structures in terms of reaction conditions, which make it suitable for high temperature conditions. The disadvantage of this approach is the time consuming procedure which involves various steps and requires preformed rigid templates. 60 Multiple templating, which combines both the soft and hard templating methodologies, is also a powerful tool for creating complex and hierarchical porous materials with greater capability and suitable for a wide range of applications, most importantly heterogeneous catalysis. 61 There are two main reasons for adopting the multi-templating approach: the first is to generate multiple porosity, i.e., the pores with different shapes and sizes, and the second is the fabrication of porous materials in desired shapes, e.g., mesoporous nanowires or mesoporous nanorods; in this approach a hard template is used to direct the shape and a soft template is used for porosity. It can be used to obtain complex porous materials with a better control of the product, but it is time consuming, needs multiple templates and it is difficult to remove the template.⁶²

Potential-controlled surfactant assembly. The potential controlled assembly within a surfactant solution is an innovative way of enabling the nanoporous metal synthesis with engineered morphology. Generally the micellar solution formed from different templates is highly sensitive to environmental stress which makes it difficult to maintain the micellar structure. 63 The electrochemical micelle assembly overcomes this limitation where the micelle formation is directed through the application of an electrical potential. In comparison with hard-templates and standard LLC soft-templates, electrochemical micelle assembly using soft-templates needs minimal steps and is therefore more cost-effective. For instance, a facile fabrication of nanoporous Pt films was reported where the strategy provides the Pt-Au alloy film with tuneable pore size ranging from 5 to 30 nm, which is effectively controlled through the molecular weight of surfactants. In this scenario, spherical micelles are formed as a result of the dissolution of nonionic surfactants in water at concentrations higher than their critical micelle concentration (CMC) values.⁶⁴ The Pt-aqua complexes are formed by the dissolution of Pt ions and coordinate with the micelle's external hydrophilic region. When an external potential is applied, the surfactant micelles containing Pt species migrate towards the working electrode. When electrons are received from the working electrode, the reduced Pt atoms in the hydrophilic region of the micelles form a stable structure with thick walls. One can see a linear increase of the film thickness with the increase in deposition time. SEM characterization showed the formation of nanopores of uniform size in the film. This shows the importance of growth control of metal nanocrystals during the fabrication of nanoporous metals in

the presence of micelles.²⁴ The interaction of micelles and dissolved metal ions influences the redox potential of metal precursors and crystal growth. Non-ionic surfactants, for example, exhibit different stabilization tendencies during the electrochemical deposition of Pd despite having similar components, and this also influences the reduction behaviour.²⁵ The strong ionic interactions between micelles and metal ions are required for system stabilization, but they are also important in increasing the barrier for metal ion reduction.⁶⁵ For instance, when the cationic surfactant cetylmethylammonium chloride (CTAC) interacts with Na₂PdC1₄, stable CTA⁺/[PdCl₄]²⁻ intermediates are formed. The electrodeposition at a high temperature (50 °C) is required to reduce the Pd precursor, which allows the formation of nanoporous films. When [PdCl₄]²⁻ is reduced, the four chloride ions released at the electrode surface cause a phase shift to rod-like micelles.⁶⁶ Under a constant potential, the cylindrical micelles align near the surface of the electrode to form a soft template for Pd film formation with perpendicular alignment of nanochannels.⁶⁷ The electrochemical micelle assembly method is adaptable and accomplished in solution, allowing for a wide range of compositions (including Pt-Au, Pt-Pd, Pt-Cu, and Pt-Ru) and shapes.62

Electrosynthesis around softly arranged colloidal assembly. The size of the pore in bimetallic alloys is often controlled by the size of the micellar hydrophobic core, so the pore size can be increased by introducing larger molecules (e.g., increasing the molecular weight of block copolymers). 68 Block copolymers with hydrophobic and hydrophilic moieties pointing inward and outward can be assembled to form stable polymeric micelles.⁶⁹ Following the formation of polymeric micelles in aqueous media, a three-electrode electrochemical arrangement can be used for the electrodeposition of nanoporous metals. Because of the formation of large size aggregates, block copolymers can precipitate at low water content. 70 By adding solvents or changing the hydrophilicity/hydrophobicity of the block copolymers, the micelle's solution stability can be maintained. The nanoporous Au film exhibits high potential in optical applications, but still the soft template based Au film fabrication remains challenging due to Au crystal growth which is difficult to control.⁷¹ In this scenario, polymeric micelle templates can be very useful. For example, by using the polystyrene-block-poly(oxyethylene) di-block copolymer (PS-b-PEO) as a pore-directing agent,72 polymeric micelles with a hydrophobic PS block as the core and a hydrophilic PEO block as the shell can be formed by regulating the solvent composition. PS-b-PEO micelles have interactions with Au species. Because of hydrogen bonding, the H₃O⁺ and AuCl₄⁻ ions released during HAuCl4 dissolution have a proclivity to react with the EO shell domains of PS-b-PEO micelles. In addition, the PS-b-PEO micelles contain free AuC1₄ ions in both solution and hydrophilic domains (Fig. 1).70 The micellar system can be charged positively or negatively or even neutral depending on the AuCl₄⁻/H₃O⁺ ratio in the EO shells. The micellar solution is slightly positive after the addition of aqueous HAuCl₄, according to zeta-potential measurements. This demonstrates that the H₃O⁺ rich micelles have a positive charge and gravitate towards the surface of the working electrode. When optimal electrochemical conditions are met, the Au nanocrystal growth is confined to the hydrophilic PEO region. The PS core forms the pores, resulting in nanoporous Au films. Under TEM, the effect of pore-directing by PS-b-PEO on nanoporous Au can be seen. Furthermore, the size and number of polymeric micelles can be easily controlled by continuously adjusting the solvent composition, which is useful in the preparation of nanoporous materials.⁷³ The nanoporous metals can be prepared by applying the electrochemical polymeric micelle assembly approach as they are difficult to synthesize by other methods.⁵⁵ The multilayered metal films of various compositions were developed using a single micellar solution (electrolyte) and perfectly programmed electrochemical plating.20

2.2 Indirect precipitation of metal hydroxides through electrogenerated hydroxide ions

Electrochemical interfacial surfactant templating. This electrochemical method has been reported for the production of nanoporous semiconductor films using surfactant self-assembly for electrodeposition.⁷⁴ It is well known that amphiphiles spontaneously aggregate at the solid-liquid interface. 75-78 Surface forces cause an increase in amphiphile interfacial concentration, resulting in the formation of surface micelles below the CMC value.⁷⁹ The nature of the substrate's surface has a significant impact on surfactant organization at the interface. Surfactant assemblies formed on a solid substrate are typically distinct from those formed in free solution. 75-78 However, the interfacial surfactant structures can be changed by modifying the chemical and electrochemical environments at the interfaces to achieve the desired structure and morphology.80 This novel strategy is typically used in combination with electrodeposition for synthesizing mesoporous metal films and alloys, where the same working electrode is used to form organic assemblies and electrochemical inorganic deposition. With the selective surfactants that provide defined hydrophilic groups for interaction with metals, the metal ions are strongly bound as counter ions on the surface of surfactant micelles, hence they form stable aggregates with organic-inorganic interfaces. When the deposition process is started by applying an electrical bias, the metal ions in the interfacial assemblies form the skeleton of the inorganic deposits, resulting in the formation of ordered inorganic nanostructures. Because this strategy only employs surfactant assemblies at solid-liquid interfaces, it enables the electrochemical formation of a wide range of inorganic nanostructures from highly dilute surfactant solutions (CMC). To improve the electrochemical interfacial surfactant templating, the impact of different chemical and electrochemical conditions on the quality, repeat distances, orientation, and homogeneity of interfacial bilayer assemblies has been investigated. These findings provide a foundation for designing and optimizing synthetic conditions for various types of metal nanostructures (for example, two-dimensional hexagonal or threedimensional cubic mesophases).81

Electro-assisted deposition through lyotropic liquid crystalline phases of surfactants. There are two major steps involved

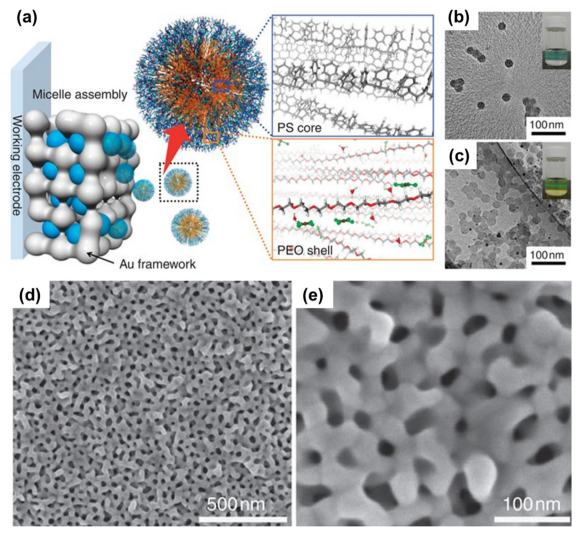


Fig. 1 (a) Schematic illustration of the mechanism involved in the formation of PS-b-PEO micelles with a gold precursor (HAuCl₄). Under a suitable composition, the PS-b-PEO block polymer reacts with Au ions to form a spherical micelle. (b) The TEM images reveal the formation of spherical micelles containing the PS₁₈₀₀₀-b-PEO₇₅₀₀ block polymer without Au ions. (c) The formation of micelles with Au ions. The black dots represent Au nanoparticles formed on the surface of micelles by electron beam irradiation. The inset image depicts the Tyndall effect. (d and e) Top-surface SEM images of the mesoporous Au film prepared at 1000 s deposition time. Reproduced with permission. 70 Copyright 2015 Springer Nature.

in the electro-assisted deposition through LLC phases. First, an LLC phase is formed over a conductive substrate, followed by metal electrodeposition using a suitable electrochemical method, and second, the surfactants are removed to yield mesoporous alloy films. To make a typical electrochemical cell with three electrodes, the counter and reference electrodes along with the working electrode are directly introduced into the as-prepared LLCs. In the metal state, a reduction of precursors is observed when an external potential is applied. The homogeneity of the LLCs is greatly responsible for homogeneously distributed pores and high quality of nanoporous structures. However, it is not easy to obtain desired mesoporous structure using LLC. This is because, LLCs are highly viscous and this high viscosity tends to agglomerate the metal ions when placed in the electrochemical cell. A few steps can be taken to avoid the problems when dealing with extremely

viscous LLCs. To begin, a diluted solution is made by combining metal precursors, a volatile solvent (ethanol, tetrahydrofuran, etc.), water, and surfactants. After the evaporation of the volatile solvent, LLCs are formed on a conductive substrate. 82,83 This method is advantageous for the subsequent preparation of various mesoporous alloy films with varying shapes and morphology.^{84,85} However, there are some drawbacks with LLC-based templating; for example, viscous LLCs reduce experimental feasibility in a three-electrode cell apparatus and limit the flexibility of the soft template process during nanoporous metal synthesis.

Sol-gel film deposition by polycondensation of hydrolyzed precursors. The sol-gel method is comprised of hydrolysis and polycondensation reactions. Typically, the sol-gel method can deposit oxyhydrate precursors and oxides that are obtained by annealing in air. This method is very straightforward and cost

effective because of no need of a vacuum system. But this method cannot be used to deposit sulfides directly. Because of the versatility of the sol-gel process, it is ideal for coating electrode surfaces with uniformly deposited metal or semimetal oxides to obtain organic-inorganic mesoporous alloy films. The method is naturally simple in that it manipulates

method cannot be used to deposit sulfides directly.86 Because of the versatility of the sol-gel process, it is ideal for coating electrode surfaces with uniformly deposited metal or semimetal oxides to obtain organic-inorganic mesoporous alloy films. The method is naturally simple in that it manipulates the fluidic character of a sol (typically made of alkoxysilane precursors for silica-based materials) for casting on a solid electrode surface, allowing gelation and drying to produce a xerogel film.87-90 The solvent evaporation steps are critical for obtaining high quality films with flat surfaces, 91 which include the ordered mesoporous nanostructures.92 On the electrode surfaces, electrolytic deposition can also be used to create sol-gel thin films. This includes immersing the electrode in a hydrolyzed sol solution and the use of an appropriate cathodic potential, which may generate hydroxyl ions locally at the electrode-solution interface. This causes an increase in pH locally and contributes to catalyze the growth of the silica film and precursor polycondensation on the surface of the electrode. 93-95 This strategy can also be used to generate organically functional silica films, 96,97 sol-gel/metal nanocomposites, 98 and conductive polymer-silica hybrids. 99 This strategy is also capable of building bio-electrocatalytic devices due to its compatibility with the encapsulation of biomolecules. 100,101 When used in conjunction with a surfactant (i.e., cetyltrimethylammonium bromide, CTAB) as a template, this method allows for the deposition of highly ordered mesoporous silica films with mesoporous channels. The orientation of the support or substrate ensures the rapid mass transportation of analytes to obtain sensitive electroanalytical devices. 102,103 Furthermore, unlike the evaporation method, this electro-assisted generation method allows uniform deposition of sol-gel layers onto the electrode surface with complex geometry and conductive patterns commonly found in Au CD-trodes, 97 macroporous electrodes, ¹⁰⁴ metal nanofibers, ¹⁰⁵ printed circuits ¹⁰⁶ or ultramicroelectrodes. 107-109 This method has been found to be suitable for producing sol-gel materials by depositing nano or micro-objects onto electrode surfaces, such as metal complexes, biomolecules (e.g., protein)¹¹⁰ or bacteria¹¹¹ that act as templates for film growth.

Electro-assisted self-assembly method (EASA). Another popular method for producing oriented mesoporous silica films is electrochemically assisted self-assembly (EASA). 112,113 This method involves CTAB as a surfactant to create nanomaterials with pore diameters ranging from 2 to 3 nm. The method entails immersing the electrode in a pre-hydrolyzed sol produced from tetraethoxysilane (TEOS) and CTAB in a moderately acidic medium (pH 3) and applying an appropriate reductive potential to generate hydroxyl ions locally at the solutionelectrode interface. The increase in local pH catalyzes the condensation of precursors 93,95 and, in the meantime, allows the formation of surfactant hemimicelles, which orients the mesoporous silica film growth perpendicular to the electrode surface via self-assembly. 114 The EASA method is useful in the preparation of functionalized mesoporous silica layers, with great promise for the elaboration of sensitive electroanalytical

3. Fabrication of mesoporous metallic alloys

3.1 Gold-silver (Au-Ag) alloy

The electrochemical fabrication of a mesoporous Au-Ag alloy can be achieved by several electrochemical methods. These distinctive methods have a significant impact on the variable properties of the synthesized alloy such as pore size and shape, pore diameter, surface area, and structure-activity relationship. The soft and hard template approach along with the electrochemical dealloying method has been extensively employed to fabricate mesoporous metals. Although a fine control over the size and structure of mesoporous metals can be achieved by the soft and hard template approach, irregular morphology of the resulting film can be a major drawback. The challenge of fabricating monodispersed mesoporous Au-Ag by the hard template approach is attributed to the faster deposition rate in mesoporous templates and formation of bulk particles due to the preferential migration to the outer layer of the mesoporous silica during the reduction process. 115 To prevent the migration of metal species outside of the mesoporous support, Fang and colleagues developed a method using a softenveloping phase transfer mechanism.116 The phase transfer mechanism is comprised of three distinct phases: solid-liquidsolution (SLS) interface where the solvent (liquid) layer acts as a barrier to block the migration of metal species to the template (solid) and cannot dissolve the used precursor molecules. The solution phase contains the reducing agent, and the final reduction of the metal precursors occurs at the interface of the solid and solution phase facilitated by the spontaneous phase transfer at the interface of the solution and liquid phase. Based on this strategy and using hexane and 1,1,3,3tetramethyldisiloxane (TMDS) as a barrier layer and reducing agent respectively on a KIT-6 hard template, monodispersed Au-Ag NPs with highly ordered uniform pores with an average diameter of ~100 nm were synthesized. The resulting Au-Ag networks exhibited higher catalytic properties in the methanol oxidation reaction (MOR) producing higher peak current up to 19.2 μ A μ g⁻¹ which is 3.1- and 24-fold higher than that of the Au-4 (nm) NPs and Au-70 NPs respectively. They also exhibited lower electro-oxidation potentials than the control Au NPs and 3- and 24-times higher mass activity and 7 and 5 times improved specific activity compared to Au-4 and Au-70 NPs respectively.

The Au–Ag alloy film can also be fabricated by using a block copolymer as a soft template that directs the pore formation and pore morphology. Such an electrochemical process of fabricating mesoporous Au–Ag alloy films has been reported by Park and colleagues. ¹¹⁷ In this work, a block copolymer templated approach was used to generate mesoporous Au–Ag

alloy films with a high surface area to volume ratio and enhanced biosensing capacity. A sacrificial soft-template, PS-b-PEO block copolymer, has been utilized as a pore directing agent.

The PS-b-PEO block polymer (dissolved in THF) forms spherical micelles upon addition of ethanol and an aqueous solution of HAuCl₄ or (and) AgNO₃ (as Au and Ag precursors, respectively). Upon the addition of the aqueous salt, the hydrophobic part of the block polymer (PS) aggregated as a core. leaving the hydrophilic EO as a shell to interact with the bimetallic species (Au and Ag ions) at the outer layer of the formed micelles. The electrodeposition of a micellar solution containing 50:50 Au³⁺:Ag⁺ at -0.5 V resulted in a homogeneously dispersed mesoporous film (Fig. 2). The as-synthesized Au-Ag alloy films were reported to contain uniformly sized pores of 12 nm throughout the alloy surface and a wider pore size of 23 nm was also found with the increase of Ag content. The resultant alloy also exhibited higher catalytic activity towards the [Fe(CN)₆]^{3-/4-} redox molecule. The homogeneous mesopores with a large surface area and the higher electrocatalytic activity of the synthesized alloy enable the construction of a highly sensitive miRNA biosensor as a transduction surface (working electrode) without requiring any enzymatic and amplification process. The magnetically isolated and purified miRNAs were absorbed onto the surface of the mesoporous Au-Ag film that contributed significantly to achieve a higher current response of 144.21 μA cm⁻² in differential pulse voltammetry (DPV) interrogation which is 18 and 2-fold higher compared to the mesoporous Ag and Au electrodes respectively. This rapid, amplification-free miRNA biosensing assay using the cost-effective Au-Ag alloy film is highly sensitive to attomolar levels (100 aM LOD) which is multiple times higher than amplification-based molecular techniques and commensurate

with the electrochemical biosensor-based detection techniques. 118,119 In terms of stability, the mesoporous Au-Ag films sustained more than 40 runs measured using cyclic voltammetry in both reaction media ([Fe(CN)₆]^{3-/4-}) and acidic media $(0.5 \text{ M H}_2\text{SO}_4).$

In another report, Lv et al. fabricated mesoporous Au-Ag nanospheres including mesoporous Au-Ag alloy, hollow mesoporous Au-Ag, and core-shell Ag-mesoporous Au. 120 A seedless, one-pot, surfactant-templating approach was utilized to fabricate the well-controlled nanostructures. A synthetic surfactant template $C_{22}H_{45}N^{+}(CH_3)_2-C_3H_6$ -SH (Cl⁻) ($C_{22}N$ -SH) along with AuCl⁴⁻ and Ag⁺ as metal precursors and N₂H₄ as the reducing agent was used to fabricate the alloys. The covalently stable intermediate generated by thiolate-metal linkages between C22N-S-Au(I) and C22N-S-Ag(I) precluded quick nucleation and high mobility of Au/Ag precursors.

3.2 Platinum-gold (Pt-Au) alloy

Although Au and Pt are referred to as the most chemically stable metals due to their resistance to oxidation reaction, synthesis of a Pt-Au bimetallic alloy is a challenging task. The imposed challenge is attributed to the different reduction kinetics of Au and Pt ions leading to phase segregation. 121-123 Massive attention has been directed towards the effective synthesis of bimetallic Pt-Au alloys due to their high catalytic properties than their pure counterparts. Moreover, the surface area of Pt-Au binary alloys largely impacts the electrocatalytic and biosensing performance since the presence of Au prevents the adsorption of poisoning intermediates such as CO by structure-induced promotional effects. 124 The challenge of fabricating a Pt-Au bimetallic alloy has been overcome by Yamauchi and co-workers, who electrochemically synthesized

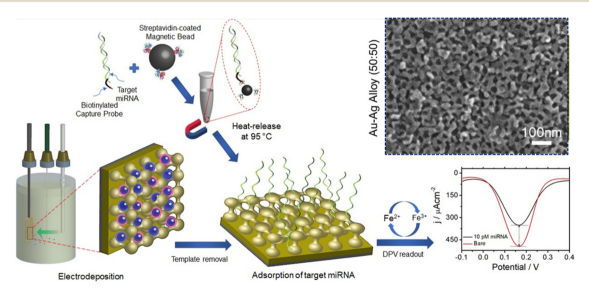


Fig. 2 Schematic diagram of mesoporous Au-Ag alloy film synthesis via the electrodeposition process. Metal precursors were deposited electrochemically onto the PS-b-PEO derived diblock-copolymer micelles followed by template removal and adsorption of magnetically isolated and purified miRNA on the Au-Aq film. Changes in the DPV current in the $[Fe(CN)_6]^{3-4-}$ system before and after the target miRNA adsorption were recorded as a part of evaluating mesoporous Au-Ag films as a single step non-enzymatic amplification free miRNA sensor. Inset: Top-surface SEM images of the mesoporous Au-Ag alloy prepared from a 50:50 precursor ratio. Reproduced with permission. 117 Copyright 2020 Royal Society of Chemistry.

a mesoporous Pt-Au alloy by employing a soft-templating strategy (Fig. 3).⁴⁰ Herein, the LLCs prepared from concentrated surfactant solution by solvent evaporation have been used as the soft-template. The use of LLCs prevented phase separation of Au and Pt ions thereby resulting in a uniform distribution despite varying deposition potentials of independent ions. They have also shown that the framework composition of the mesoporous structure can be readily controlled by tuning the ratio of the precursor solution. Shifting the Au content from low to high yielded distorted mesoporous structures (Pt₆₂Au₃₈) compared to the well-ordered 2D hexagonal mesostructures (Pt₉₀Au₁₀ and Pt₈₀Au₂₀) which in turn reduced the electrochemically active surface area. In SEM analysis, pores were reported to be distributed over the entire surface of mesoporous alloy films in a honeycomb arrangement with a pore diameter of around 3 nm. When compared to nonporous Pt sheets and mesoporous Pt, the resulting mesoporous alloy performed admirably as a catalytic electrode in electrochemical methanol oxidation reactions, exhibiting substantial current density for a longer period. This durable feature was achieved due to the prevention of electrode deactivation by poisoning in the presence of Au which is absent in nonporous and mesoporous Pt. 125

Another surfactant based templating strategy using poly-(ethylene glycol)-poly(propylene glycol)-poly(ethylene glycol) (P123) as a template was reported for the fabrication of Pt-Au mesoporous alloy films by electrodeposition on indium-tin oxide (ITO) coated glass. 125 The P123 template contributed to the surface morphology by affecting the growth of particles. The fabricated Pt-Au alloy films also exhibited higher catalytic activity in the methanol oxidation reaction and induced a peak area increase and peak shift to lower potential in the I-V curve (at 50 mV s⁻¹ scan rate) compared to the fabricated nonporous and mesoporous Pt electrode. In the performance characteristics test of the synthesized film in the methanol oxidation reaction, the Pt-Au film displayed a higher current density for a longer period of time compared to the nonporous Pt and mesoporous Pt indicating the alloy's significant stability.

In 2013, Cuiling Li and colleagues reported another approach of electrochemical synthesis of a mesoporous Pt-Au alloy film by a soft-templating method.²² In their work, they were able to fabricate a mesoporous Pt-Au alloy by electrochemical deposition on a Au-coated Si substrate. Rather than applying constant potential deposition, a square-wave potential program was applied in the process. Application of a squarewave potential facilitated the uniform reduction and constant concentration of metallic ions on the substrate surface resulting in the formation of a Pt-Au alloy with a pore size of approximately 7-8 nm and without any phase separation. The square-wave potential also permitted the retention of a constant mesoporous structure with up to 50% atomic content of Au whereas a distorted mesoporous structure in the Pt₆₂Au₃₈ alloy was noticed in the absence of a square-wave potential. The fabricated Pt-Au alloy films were found to be a highly effective glucose sensor owing to their high catalytic activity towards glucose oxidation. This is due to the synergy of the large surface area of the Pt-Au alloy and mesoporous structures. As an amperometric glucose sensor, the fabricated mesoporous Pt51Au49 alloy was reported to be highly sensitive (352 μ A cm⁻² mM⁻¹) with a lower detection limit (LOD) of 6 μM and a detection range of 6 μM - 11 mM which is within the normal physiological level of glucose (8-11 mM). 126,127

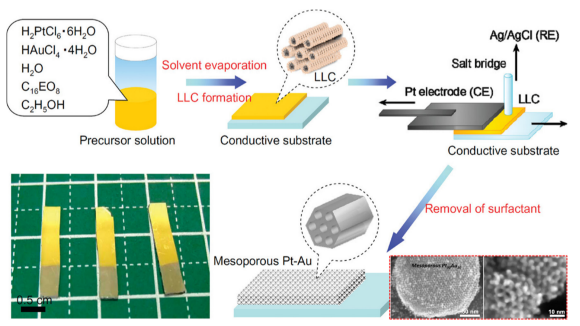


Fig. 3 Illustration of the experimental steps involved in the fabrication of the mesoporous Pt-Au alloy film. The highly ordered two-dimensional LLC template was prepared on a conductive surface by a drop-coating method from a surfactant solution containing ethanol, water, nonionic surfactant, and the corresponding metal species. The resulting LLC film comprising Pt and Au species was used to perform metal electrodeposition utilizing a galvanostat and a Pt substrate as the counter electrode and Ag/AgCl as the reference electrode, followed by removal of surfactants and undeposited metal species. Inset: SEM images of mesoporous Pt₉₀Au₁₀. Reproduced with permission.⁴⁰ Copyright 2012 American Chemical Society.

Moreover, after their catalytic activity, the native state of the Pt-Au electrodes can be achieved with a simple rinse with distilled water and the electrode structure can sustain at least 10 cycles of glucose detection. Electrochemically fabricated mesoporous Pt-Au binary alloy films (Pt51Au49) featuring high sensitivity, low LOD, wider detection range, high fabrication reproducibility, and anti-poisoning effects enable the application of the Pt-Au alloy as a non-enzymatic glucose sensor for a wide range of clinical purposes.

3.3 Gold-copper (Au-Cu) alloy

Highly viscous and concentrated surfactants used in a typical liquid crystal-based synthesis of mesoporous alloys interfere with the pore size tuning of the fabricating alloy.³⁴ To get around the problem, researchers have employed a combination of electrochemical deposition, polymeric micelle assembly, and easy solvent extraction to create a mesoporous Au-Cu alloy with variable composition. Nugraha et al. have reported the synthesis of Au-Cu mesoporous alloy films by electrodeposition on Au-Si substrates using spherical polymeric micelles as the electrolyte solution.²³ In this strategy, the PS-b-PEO polymeric micelle was used as a sacrificial soft-template. Fine control over the architecture of mesopores was achieved by tuning the compositional ratio of precursor species (Au and Cu) present in the electrolyte solution. Uniform distribution of the mesopores was reported in Au₉₄Cu₆, Au₈₇Cu₁₃, Au₇₈Cu₂₂, Au₄₉Cu₅₁, and Au₄₁Cu₅₉ alloy samples with an average pore diameter of 28 nm (Fig. 4). Interestingly, it was found that the final structures of the alloy mimic the soft template used as a support. Au₉₄Cu₆ and Au₈₇Cu₁₃ alloys which were low in Cu content appeared to replicate the cage-like mesopores present in the parent micellar template. In contrast, the higher Cu content in Au₇₈Cu₂₂ and Au₄₉Cu₅₁ alloys mediated the fusion of mesopores resulting in one-dimensional vertically aligned mesochannels. These vertical channels provide lesser diffusion resistance which in turn facilitates mass transfer within the mesoporous channels. Electrochemically synthesized Au-Cu

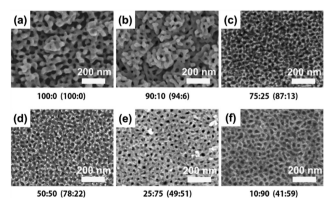


Fig. 4 The top-surface SEM images of Au-Cu alloy films with different compositions of both Au and Cu, fabricated by using the PS-b-PEO polymeric micelle. The presence of mesopores in the Au-Cu alloy samples confirms the formation of mesoporous films. Reproduced permission.²³ Copyright 2018 American Chemical Society.

bimetallic mesoporous alloy films were also found to be highly active in the electro-oxidation of glucose. In general, the performance of a non-enzymatic glucose sensor heavily depends upon two critical parameters: electroconductivity and the large surface area of electrodes that are crucial for electro-oxidation of glucose.23 All of the Au-Cu alloys possessed a high volumenormalized electrochemically active Au surface area (ECSA) compared to the Au films. Moreover, the higher current response and more positive potential of bimetallic Au-Cu mesoporous alloys compared to the mesoporous Au films in the electro-oxidation of glucose characterize the alloys to be more catalytically active. The mesoporous Au₄₉Cu₅₁ film was also found to be highly sensitive (643.6 μA cm⁻² mM⁻¹) than the nonporous Au₄₉Cu₅₁ film and mesoporous Au film over a broad range of glucose concentration (0.01 – 19 mM) with a LOD of 1.5 μM. To date, there are only two Au- and Cu-based nanomaterial-based glucose sensors available that possess higher sensitivity than the mesoporous Au-Cu film.^{128,129}

3.4 Gold-nickel (Au-Ni) alloy

Application of electrochemical deposition to synthesize mesoporous alloy films can be extended further to fabricate Au-Ni bimetallic alloy films with mesoporous morphology. Such an approach was reported by Nugraha and colleagues where a block-copolymer template was utilized to synthesize mesoporous Au-Ni alloy films with different compositions. 26 In this soft-template mediated electrodeposition, Au and Ni precursors were reduced electrochemically in the presence of PS-b-PEO diblock copolymer micelles. The resultant Au-Ni films consist of mesopores distributed uniformly throughout the entire surface with an average pore size of 28 nm. By adjusting the Au: Ni ratio in the precursor solution, several Au-Ni alloy films with varying compositions (Au₁₀₀, Au₈₁Ni₁₉, Au₇₆Ni₂₄, Au₆₀Ni₄₀) were produced. A negative correlation between alloy thickness and Ni²⁺ content in electrolyte solution was observed. The welldefined mesopores are generated throughout the film, from the outermost to the innermost parts (Fig. 5). By enhancing accessibility to active sites and lowering mass and ion transport resistance, such a mesoporous architecture can be of considerable benefits for electrocatalytic applications, allowing for an optimum contact area between the catalyst and the analyte. Mesoporous Au_{100} , $Au_{81}Ni_{19}$, $Au_{76}Ni_{24}$, and $Au_{60}Ni_{40}$ had a Aumass-normalized ECSA-Au of 113.9, 38.2, 94.2, and 53.4 cm² mg^{-1} _Au, respectively. In the cyclic voltammetry (CV) test to measure the electrocatalytic performance toward the methanol oxidation reaction (MOR), mesoporous alloy films exhibited two characteristic anodic peaks during positive and negativegoing scans; the former is due to the oxidation of methanol whereas the latter is linked to the removal of the leftover carbonaceous intermediates. 130,131 The catalytic current density of alloy compositions increased gradually with the increase of Ni content until reaching the optimum for Au₇₆Ni₂₄ (410 mA mg $^{-1}$ _Au) and then declined. This obtained current density is the highest among other reported catalysts used for MOR. 132-135 In durability check of the binary Au-Ni alloy for MOR by chronoamperometry, Au₇₆Ni₂₄ exhibited stable current

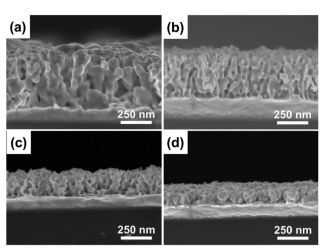
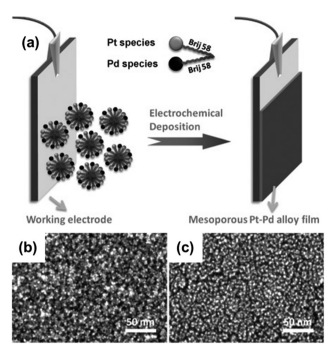


Fig. 5 The cross-sectional SEM images of mesoporous bimetallic Au-Ni films obtained from precursor solutions using PS-b-PEO diblock copolymer micelles containing different molar ratios of Au and Ni (a) 100:0, (b) 75:25, (c) 50:50, and (d) 25:75 at an applied potential of -0.7 V and temperature of 40 °C. Reproduced with permission.²⁶ Copyright 2019 Flsevier Ltd

density and the lowest deterioration rate of current density for 1200 seconds among other compositions owing to the prevention of poisoning by intermediate carbonaceous species. 136 The simple preparation reinforced by excellent electrocatalytic capability and stability underscores the practical implementation of the mesoporous Au₇₆Ni₂₄ bimetallic alloy film in direct methanol fuel cells (DMFCs).

3.5 Platinum-based (Pt-Pd/Ru/Cu) alloy

Unlike other bimetallic alloys, platinum-palladium alloys are in extensive use as catalysts in several application fields including fuel-cell technology, hydrogenation reactions, hydrocracking, hydrotreatment, hydro-isomerization, and so on. 137 But the synthesis of Pt-Pd mesoporous alloys requires a lot of complex processes with poor control over the framework composition. Wang and Yamauchi demonstrated in 2012 that electrochemical plating in an aqueous surfactant solution may be used to make mesoporous Pt-Pd alloy films (Fig. 6).24 The aqueous electroplating solution contained K₂PtCl₄ and Na₂PdCl₄ along with a non-ionic surfactant Brij 58. Brij 58 was taken at a concentration of 1.0 wt%, which is above its CMC. By changing the compositional ratio of precursor species, compositional control over the mesopores of the synthesized alloy (Pt₉₃Pd₇, Pt₈₂Pd₁₈, Pt₆₉Pd₃₁, and Pt₅₇Pd₄₃) was achieved. The resulting film contained mesopores distributed uniformly throughout the surface with a pore size of 7 nm and a wall thickness of 3 nm. From the SEM image, it was evident that the absence of the platinum precursor induced the palladium to develop as a branch without forming any mesopore whereas an increased content of palladium caused visible cracks and an uneven alloy surface. The same effect was also reflected upon the alloys' catalytic performance in the methanol oxidation reaction where Pt₆₉Pd₃₁ and Pt₅₇Pd₄₃ exhibited lower catalytic activity than mesoporous platinum and Pt93Pd7 and Pt82Pd18 alloy films.



(a) Pictorial representation of the mesoporous bimetallic Pt-Pd alloy film synthesis by an electrochemical plating approach where Pt and Pd metal species are electrodeposited into the working electrode from a precursor solution containing K₂PtCl₄, Na₂PdCl₄, and non-ionic aqueous surfactant Brij 58. High-resolution SEM images of mesoporous Pt (b) and Pt57-Pd43 alloy films (c). (b) Reproduced with permission.²⁴ Copyright 2012 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim.

Pt93Pd7 and Pt82Pd18 achieved increased catalytic activity of 30% and 60% respectively than the mesoporous platinum film. This enhanced performance is due to the high palladium content which reduces the energy for methanol adsorption and facilitates dehydrogenation of methanol to adsorbed CO.^{138,139}

In a regular platinum catalyst, CO can trigger the poisoning of the platinum surface. But the synergy between platinum and palladium in the Pt-Pd alloy film alleviates CO poisoning. Moreover, the presence of platinum nanoparticles in the wall of mesopores also boosts up the methanol oxidation reaction.⁶³ Alongside catalytic performance, the Pt-Pd film also possessed significant stability in CV analysis as it achieved the highest current density for 2000 seconds compared to the mesoporous Pt and Pt catalyst. Similar to the Pt-Pd alloy, mesoporous Pt-Ru alloy film synthesis of different compositions has been reported by Wang and colleagues.⁶⁶ In their work, they have used Brij58 as a non-ionic surfactant solution and K2PtCl4 and RuCl₃ as metal precursors to perform electroplating on gold (Au) surfaces (Fig. 7). Different compositions of Pt and Ru in the precursor solution (Pt^{2+}/Ru^{3+}) such as 100:0,75:25,50:50,and 25:75 resulted in alloys of Pt100, Pt95Ru5, Pt93Ru7, and Pt₈₇Ru₁₃ respectively. Surprisingly, a substantial increase of Ru species in the precursor solution did not correspond to the amount deposited on the films (50% in the precursor to mere 7% in the product). This unique result could be attributed to the lower reduction tendency of Ru³⁺ ions than [PtCl₄]²⁻ in the

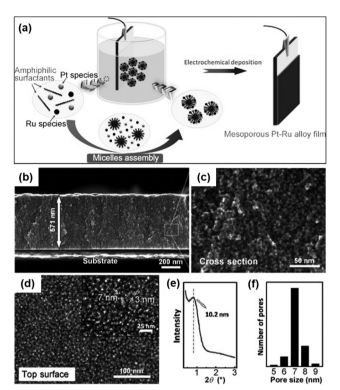


Fig. 7 (a) Fabrication process of the mesoporous Pt-Ru alloy film by using electrochemical reduction using Brij58 as an amphiphilic surfactant and K₂PtCl₄ and RuCl₃ as Pt and Ru precursors respectively. (b) Crosssectional SEM image at low magnification. (c) Corresponding highly magnified SEM image. (d) Top-surface SEM image. Inset: Highly magnified image of the top surface. (e) Low-angle XRD profile. (f) Pore-size distribution curve in the film. Reproduced with permission.⁶⁶ Copyright 2012 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim.

reduction process during electrodeposition leading to the preference of [PtCl₄]²⁻ ions over Ru³⁺ ions. 140 The as-synthesized alloy films possessed uniformly distributed mesopores with a diameter of 7 nm along with inter-connected nanoparticles in the pore wall. These connected nanoparticles contributed greatly to attain a concave surface offering numerous rich edges and corner atoms. 141 In the methanol oxidation reaction, the mesoporous Pt95Ru5 alloy exhibited increased catalytic activity compared to the Pt nanoparticles, mesoporous Pt films, and the commercially available Pt catalyst. The chronoamperometric curve of the Pt-Ru alloy against the mesoporous Pt and Pt catalyst in the methanol dehydrogenation reaction showed considerably higher current density indicating good stability of the alloy film.

Another study focusing on the electrochemical method to synthesize mesoporous Pt-Cu alloy films has been reported by Li and Yamauchi in 2014.²⁵ Similar to the Pt-Pd and Pt-Ru, mesoporous Pt-Cu films were electrochemically deposited on a gold-coated silicon substrate (as a working electrode) at a reduction potential of -0.2 V in an electrolyte solution containing metal precursors and non-ionic surfactant Brij58. A linear relationship between the metal content in the precursor solution and the resulting films of various Pt and Cu compositions was reflected. However, a shift of surface morphology

from flat to a cauliflower-like structure with the increase of Cu content was reported from SEM images. From the synthesized alloy films of different compositions (Pt₁₀₀, Pt₈₆Cu₁₄, Pt₇₃Cu₂₇, Pt₅₇Cu₄₃, Pt₃₉Cu₆₁ and Pt₃₀Cu₇₀) the thickness of Pt₈₆Cu₁₄ and Pt₅₇Cu₄₃ was reported to be similar around 550 nm. In the catalytic performance test towards the methanol oxidation reaction (MOR), Pt₅₇Cu₄₃ generated the highest current density of 1.02 mA cm⁻² Pt compared to 0.98 mA cm⁻² Pt and 0.96 mA cm⁻² Pt for Pt₈₆Cu₁₄ and Pt₁₀₀ respectively. Moreover, a negative shift of the onset potential and peak potential in mesoporous Pt-Cu films was evident compared to the mesoporous Pt film. A comparatively higher current density of Pt₅₇Cu₄₃ than Pt₈₆Cu₁₄ despite sharing other properties was due to the fine balance between Pt and Cu content where Cu scavenges poisonous intermediates generated on the Pt electrodes suppressing CO poisoning. 142 However, the mesoporous Pt-Au alloy possess relatively less stability (long-term) towards the MOR than that of mesoporous Pt. This is probably due to the less stability of individual Cu for MOR, that also shown by Pt NPs catalyst for MOR. 143

Rhodium-nickel (Rh-Ni) alloy

Nanomaterials based on rhodium (Rh) have been used substantially in various applied fields such as fuel cell technology, synthesis of refined chemicals, etc., owing to their high electrocatalytic activity. 144,145 However, several associated factors such as complexity in the construction of high-surface-area structures due to high surface energy, less control over the final product, and high cost of Rh metal render the traditional wet chemical synthesis of Rh nanoparticles an inefficient process. 146 To overcome the issue, Kani and colleagues undertook an alloying approach of Rh with inexpensive transition metal Ni to fabricate mesoporous RhNi alloy films via an electrochemical method (Fig. 8).21 Metal precursors of Rh and Ni were co-deposited on Au-coated Si substrates by a selfassembled PS-b-PEO block copolymer micelle. To identify the optimal electrochemical deposition potential for RhNi films, linear sweep voltammetry (LSV) analysis was performed, and Rh deposition started at -0.2 V irrespective of the presence of Ni precursors in solutions. Since Ni has a much lower redox potential (E^0) than Rh (-0.26 V and 0.76 V respectively), it required a much higher energy and low potential (-1.0 V)and a lowered pH of 2.4. But at low potential, fast deposition of Rh occurs that hinders incorporation into the micelles. Hence the co-deposition of Ni on the Rh layers occurs to generate mesoporous RhNi alloys at -0.7 V deposition potential. Though the Ni content in the final alloy materials corresponds linearly to the Ni concentration in the precursor solution, the final alloy films did not exhibit more than 50% of Ni content, supporting the co-deposition phenomena. By tuning the composition in precursor solutions, Rh₁₀₀, Rh₇₆Ni₂₄, Rh₅₈Ni₄₂, and Rh₄₉Ni₅₁ films with uniformly distributed pore diameters of 17.0, 16.5, 16.0, and 16.6 nm respectively were produced.

The high-angle annular dark-field scanning TEM (HAADF-STEM) revealed that the pore walls consisted an array of tiny

Micelles assembly (a) Template removal PEO-b-PMMA Polymer/metal ion Ni²⁴ composite micelle Mesoporous Rh, Ni, 100, , flim Electrodeposition

Fig. 8 (a) Schematic representation of the electrochemical fabrication of mesoporous Rh-Ni alloy films. A PEO-b-PMMA block copolymer template was utilized to electrochemically deposit Rh and Ni metal species on a gold (Au)-coated silicon (Si) substrate followed by the template removal step to generate a mesoporous bimetallic Rh-Ni alloy film. Top-surface SEM images of mesoporous (b) $Rh_{76}Ni_{24}$, (c) $Rh_{58}Ni_{42}$, and (d) $Rh_{49}Ni_{51}$ films deposited at -0.7 V for 270, 240, and 180 s, respectively. Reproduced with permission. 21 Copyright 2021 Royal Society of Chemistry.

nanomaterials contributing to the increase of surface area. Formation of alloy crystals with Ni and Rh was also proved by peak shift towards larger angles in XRD with the increase of Ni concentration. The catalytic performance of the synthesized alloys was evaluated by alkali-mediated hydrogen evolution reaction (HER), which serves as a model reaction for the performance test. The ECSA_Rh of Rh₁₀₀, Rh₇₆Ni₂₄, Rh₅₈Ni₄₂, and Rh₄₉Ni₅₁ was reported as 39.9, 40.6, 44.0, and 34.3 m² g⁻¹ respectively. RhNi films also exhibited a positively shifted polarization curve compared to the Ni₁₀₀ film (> -100 mV νs . -276 mV) indicating higher production of hydrogen. The performance in HER was reported to increase with the increase of Ni concentration in the alloy, supported by current densities of -72, -65, and -59 mV (a) 10 mA cm⁻² for Rh₇₆Ni₂₄, Rh₅₈Ni₄₂ and Rh₄₉Ni₅₁ alloys respectively. Tafel slopes, an indicative measure for the HER performance (the lower the better), also indicate the high-performance rating of the mesoporous $Rh_{49}Ni_{51}$ alloy film (67 mV dec⁻¹ compared to 69, 74, 91, and 100 mV dec⁻¹ for Rh₅₈Ni₄₂, Rh₇₆Ni₂₄, Rh₁₀₀, and Ni₁₀₀) among other alloys of varying compositions. 147-149 In chronopotentiometry examination of the Rh-Ni alloy film for 10 straight hours, the catalytic activity decreased over time compared to the pure Rh film. Although pore morphology and Rh-Ni composition remained unchanged, the properties of catalytically active sites were reported to be affected. It is due to the common characteristics of Ni and its alloys to produce Ni hydrides (NiH_x) on the electrode surface during prolonged activity thus lowering the performance in HER. 150

3.7 Mesoporous chalcogenide semiconductors (Cd-Se and Cd-Te)

In recent years, a new avenue of research has opened up focusing on mesoporous semiconductors, a new class of materials with potential applications in optoelectronics and solar cells. The adjustable morphological aspects of mesoporous semiconductors (particle size, pore size, etc.) that directly influence the light scattering pattern inside the nanoporous structures make them more promising. Though there are several methods currently available for synthesizing chalcogenide semiconductors, the reports about mesoporous CdX (X = chalcogenide) are very limited. In 2020, Nagaura and colleagues reported for the first time an electrochemical method for fabricating CdSe and CdTe films (Fig. 9).151 A soft-template directed electrodeposition was performed using PS-b-PEO block copolymer micelles. The pore size of the fabricated films was found to be substantially dependent on the molecular weight of the block copolymer, with pore diameters of 9-9.5 nm and 17.7-18.0 nm for PS(5000)-b-PEO(2000) and PS(10 000)-b-PEO(4100) respectively. From the XRD pattern, it was proved that the CdSe and CdTe films posed as a hexagonal/cubic mixture and cubic structure respectively. By using this as-prepared mesoporous CdSe film and non-porous CdSe film as the photo-absorber, they also developed highly sensitive photodetectors. The photo-to-dark current ration (PDCR) which measures the photodetection sensitivity of the non-porous CdSe film was reported as 0.059 at 0.05 mW cm⁻² and the optical sensitivity (S) was 1.18 mW⁻¹ cm² compared to 10 and S = 204 for the mesoporous CdSe based photodetector. Massive improvements

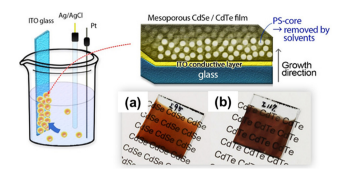


Fig. 9 Schematic representation of the fabrication of mesoporous CdTe and CdSe films, and their morphological structure: (a) CdSe and (b) CdTe film. The films were grown onto an ITO glass substrate using PS-b-PEO block copolymers. Reproduced with permission. 151 Copyright 2020 Wilev-VCH GmbH.

of the mesoporous CdSe based photosensor are due to two important factors. Firstly, the low dark current of the mesoporous CdSe than that of the non-porous samples, and smaller grain and pore size (~ 9 nm) than the mean free path of electrons of n-type CdSe (≈20 nm) result in a larger PDCR value. 151 Secondly, numerous electrical semiconducting nanojunctions inside of mesoporous architecture can be directly exposed to the light as light can penetrate and scatter through the nanopore networks, substantially enhances the quantum efficiency of the photosensors. 152,153

3.8 Mesoporous trimetallic alloys

Alloying of metals by electrochemical fabrication methods with tunable compositions can also be extended to trimetallic alloys. Along with lowering the production cost, electrochemically

synthesized trimetallic alloys offer enhanced catalytic performance with a high degree of reaction stability. 154-156 Electrochemical fabrication methods also offer a multitude of options to optimize alloy composition to adjust the catalytic performance of the resultant ternary alloys. Such a study has been reported by Nugraha et al. in 2018, where the group synthesized mesoporous AuCuNi films by electrochemical deposition (Fig. 10). 157 The co-electrodeposition of Au, Cu, and Ni (HAuCl₄, CuSO₄, and NiCl₂ as precursors respectively) on a conductive Au-coated Si surface using PS-b-PEO polymeric micelles as a sacrificial template resulted in mesoporous AuCuNi films having a uniform distribution of mesopores of 28 nm throughout the entire surface. The synthesized films were also reported to exhibit a strong dependency on the metal precursors present in the micellar (electrolyte) solution. By adjusting the precursor solution's composition, mesoporous Au₃₇Cu₄₈Ni₁₅, Au₄₁Cu₄₆Ni₁₃, and Au₆₀Cu₃₅Ni₅ films were obtained. Interestingly, SEM, TEM, EDS, and XRD analysis of Au₆₀Cu₃₅Ni₅ films revealed that the mesoporous structures were not confined only to the surface area of the film, rather they expanded across the entire cross-sectional area. CV analysis of the synthesized alloy films showed an ECSA of 43.2, 45.8, and 18.1 $\text{m}^2\text{ g}^{-1}$ for Au₃₇Cu₄₈Ni₁₅, Au₄₁Cu₄₆Ni₁₃, and Au₆₀Cu₃₅Ni₅ respectively indicating a larger contact area and easily accessible pores available for welcoming guest species to undergo electrocatalytic reaction.

Electrochemically fabricated mesoporous AuCuNi films have displayed intensified activity in the electrocatalysis of glucose compared to the pure (non-porous) Au film. 72 Mesoporous Au₆₀Cu₃₅Ni₅ was able to detect glucose at concentrations ranging from 0.1 mM to 8.0 mM with a detection sensitivity of 1823 µA mM⁻¹ cm⁻² and a detection limit of

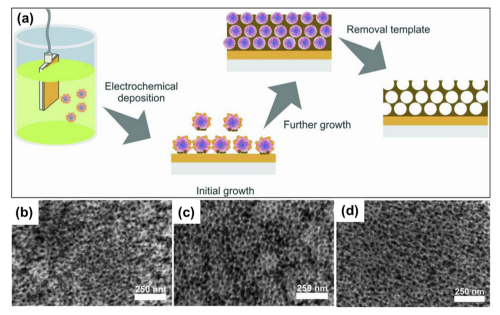


Fig. 10 Mesoporous trimetallic alloy synthesis process by electrodeposition involving the polymeric micelle assembly strategy. The ternary AuCuNi alloy with well-organized mesopores was synthesized by co-electrodeposition of Au, Cu, and Ni precursors on an Au-coated Si surface in the presence of PS-b-PEO polymeric micelles. SEM images of the mesoporous AuCuNi films synthesized from a Au: Cu: Ni precursor ratio of 3:1:1 at an applied potential of (b) -0.70 V, (c) -0.80 V, and (d) -0.90 V (vs. Ag/AgCI) at 40 °C for 1000 s. Reproduced with permission. ¹⁵⁷ Copyright 2018 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim.

0.82 µM, which is relatively higher than most of the nonenzymatic glucose sensing electrodes. The fabricated ternary alloy film is also reported to be immune to the naturally occurring interfering agents such as maltose, uric acid, fructose, and ascorbic acid in glucose sensing warranting its high selectivity. Moreover, in the methanol oxidation reaction, all of the AuCuNi alloys exhibited increased peak current density where the highest current density was achieved with Au₄₁Cu₄₆Ni₁₃ (3.8 mA cm⁻²), which is also notably higher than that achieved with Pt-based electrocatalysts. 158,159 The corresponding ternary alloys also presented a great deal of stability in retaining their activity in glucose and methanol oxidation reactions by minimizing the loss of ECSA in respective reactions. The ECSA of the ternary alloy decreased merely 19% after 6500 seconds of electro-oxidation of glucose and the alloy showed a high catalytic activity in the methanol oxidation reaction evaluated by the amperometric response for 1200 seconds. Another insightful study by Nugraha and colleagues undertook an exploratory approach by Bayesian optimization to find the ternary PtPdAu alloy with optimum catalytic performance in the methanol oxidation reaction. 42 This exploratory search curtailed the need for time-consuming and exhaustive experimentation to filter out the chemical composition of the most efficient mesoporous PtPdAu alloys. To put the numbers in perspective, possible compositions of individual metal species to fabricate trimetallic alloys can be 5151 which makes it very challenging to determine the proper composition for obtaining trimetallic alloys with highest electrocatalytic activity. Mesoporous trimetallic PtPdAu alloys with varying compositions were electrochemically synthesized by the micelle assembly approach using the Pluronic P123 EO₂₀-PO₇₀-EO₂₀ triblock polymer and K₂PtCl₄, Na₂PdCl₄, and HAuCl₄ as metal precursors. Structures with well-defined and well-distributed mesopores with 9 nm pore size devoid of any cracks were reported, which mimic the mesopores of the micelle template. The synthesized alloys also manifested a higher catalytic activity in the methanol oxidation reaction generating an oxidation peak in the forward scan associated with methanol oxidation and a peak in the backward scan associated with removal of reaction intermediates (CO for example) generated in the forward scan. 160-162 The obtained current responses were reported to be higher than those of the bimetallic alloy of PtPd and PtAu supporting the synergistic effects of the third metal species in electrocatalytic performance. 162,163 Of the various compositional ratio of each metal species in the precursor solution, the one with the lowest Au content (Pt:Pd:Au ratio of 59:40:1) generated the best current density in methanol oxidation, which was also the composition derived from the Bayesian optimization. This might be attributed to the change of surface adsorption of Pt by Au leading to better CO removal and catalytic performance. 164,165

4. Application of mesoporous metallic alloy films

Mesoporous metallic alloy films provide significant promise for advanced technology development in response to the most pressing global concerns (*e.g.*, energy generation, storage, water decontamination, wettability, automated device fabrication, photosensing, biosensing and nanomedicine).^{34,166–168} These materials are excellent candidates for developing advanced technological applications which range from hydrogen production to fuel cells and sensors, not only because they possess special physiochemical properties, but also because of the high flexibility of their synthesis.

4.1. Hydrogen evolution reaction (HER)

Fuel cell development is accelerating in response to current environmental concerns and the growing demand for more efficient power sources. 169 The fuel cell, which transforms the chemical energy of a fuel directly to electrical energy, has numerous advantages over traditional combustion-based technologies. 170,171 Hydrogen has emerged as a possible successor for fossil fuels, as a clean and sustainable energy with the highest energy density. When used as a fuel in a fuel cell, it not only converts energy efficiently, but also creates no pollution because it only emits water as a byproduct. However, yet it is very challenging to generate a large quantity of hydrogen that can serve commercial purpose. Water splitting electrolysis is a viable technique for achieving efficient hydrogen synthesis in terms of energy conversion and storage, in which catalysis or electrocatalysis plays a vital role. The hydrogen evolution reaction (HER) produces hydrogen from water electrolysis. Owing to the excellent intrinsic activity, Pt-group metals such as Pt, Pd, and Ru have attracted increased study interest in recent decades as excellent HER catalysts.172 However, expensive synthesis procedures and scarcity of these metals limit their widespread use for HER.¹⁷³ As a result, developing effective, robust, and inexpensive catalysts or electrocatalysts is a must if meaningful electrocatalytic hydrogen production from water splitting is to be achieved. Mesoporous Rh-Ni alloy films fabricated by an alloying approach of Rh with inexpensive transition metal Ni via an electrochemical method resulted in improved catalytic performance as an electrocatalyst in HER.21 The performance of the alloy in HER improves with the increase of Ni concentration in the alloy, and the mesoporous Rh₄₉Ni₅₁ alloy film is reported to be the best supported by a current density of -59 mV @ 10 mA cm⁻² (Fig. 11). Tafel slopes, another indicative measure for the HER performance (the lower the better), also indicate the high-performance rating of the mesoporous Rh49Ni51 alloy film. The alloy's mesoporous structure and favourable synergistic electronic characteristics are credited with the improved HER performance.²¹ Again mesoporous Pt with $\Delta GH \approx 0$ is well known as the bestperforming catalyst for the HER, which requires a negligible over-potential to achieve high reaction rates in acidic solutions. 174,175 However, its functionality as the HER catalyst in practical use is limited by the high cost of this metal. Mesoporous Ni-rich Ni-Pt thin films were produced by alloying Pt with the transition (relatively inexpensive) metal Ni which significantly reduced the cost of the whole catalyst without compromising catalytic activity. Also, the secondary metal heteroatoms optimized the d-band center and Δ GH of Pt which

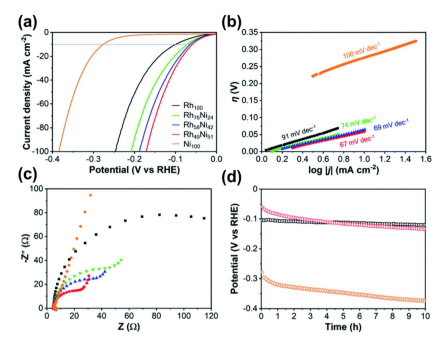


Fig. 11 Demonstration of the alkali-mediated HER performance of the Rh-Ni alloy in 1.0 M KOH solution. (a) The polarization curves of mesoporous Rh_{100} , $Rh_{76}Ni_{24}$, $Rh_{58}Ni_{42}$, $Rh_{49}Ni_{51}$, and Ni_{100} films recorded at a scan rate of 2 mV s⁻¹. (b) Tafel plots and (c) EIS of the corresponding samples. (d) The HER stability of mesoporous Rh100 (black), Rh49Ni51 (red), and Ni100 (orange) films demonstrated using chronopotentiometric V-t graphs obtained at a current density of 10 mA cm². Reproduced with permission.²¹ Copyright 2021 Royal Society of Chemistry.

resulted in enhanced performance and excellent stability of the entire bimetallic system. The HER activity is found to be increased by almost one order of magnitude compared to Pt/C.176

4.2. Methanol oxidation reaction (MOR)

When compared to hydrogen, methanol is a very appealing fuel because of its availability, low cost, and high energy density, as well as its ease of storage and transportation. 177-179 Because of their portability and promise of high energy density, DMFCs have sparked a lot of scientific interest. 177,178,180 Without reforming the fuel into hydrogen, DMFCs may generate power from methanol and water at low temperatures. 181 While DMFCs are expected to be one of the future energy sources in a hypothetical carbon-neutral economy, there is still a significant gap in the development of a suitable electrocatalyst, which necessitates further research. 179 Though mesoporous Pt with a high surface area demonstrates significant activity in the electrocatalytic oxidation of methanol, which is a halfreaction of a DMFC, the adsorbed CO molecules formed as a reaction intermediate hamper the reaction by obstructing the active sites of Pt. 182 As a result, tremendous work has gone into preventing CO poisoning. Yamauchi and colleagues undertook an alloying approach of Pt with Au to fabricate a Pt-Au mesoporous film which tends to exhibit an excellent electrocatalytic performance for methanol oxidation reactions. The presence of Au suppresses the adsorption of CO onto the Pt surface, to alleviate its poisoning effects (Fig. 12). 40 In terms of electrochemical stability toward methanol oxidation, the stability of mesoporous Pt-Au alloy films exceeds that of nonporous Pt and mesoporous Pt films as well as the commercial Pt black powders.40 Similar to the mesoporous Pt-Au alloy film, other Pt-based mesoporous alloy films such as Pt-Ru and Pt-Cu films also showed strong potential for use as an electrocatalyst in direct methanol fuel cells.^{25,66} Likewise, the excellent electrocatalytic capability and stability of the mesoporous Au-Ni bimetallic alloy film fabricated by electrochemical deposition underscores the practical implementation in DMFCs.²⁶ Furthermore, compared to binary alloys, mesoporous trimetallic alloy catalysts such as mesoporous Au-Cu-Ni films and Pt-Pd-Au films have been observed to exhibit greater overall catalytic activity. 42,157,183

4.3. Glucose sensor

Glucose sensors have received increased attention in the medical use for blood glucose measurement and monitoring over the last few decades. The accurate monitoring and careful control of the amount of glucose in human blood are essential. 184,185 However, the last few decades have experienced an obvious rise in diabetes cases worldwide. This has resulted in a surge in demand for non-enzymatic amperometric glucose sensors that are extremely efficient, stable, selective and sensitive. Noble metals, such as Pt and Au, have been identified as the most promising candidates for glucose electrooxidation. 23,186,187 These metals as electrodes, on the other hand, have limited sensitivity and poor selectivity, and are poisoned by reaction intermediates. Some of these flaws were addressed by combining various noble metals. Mesoporous Pt-Au alloy films, for example, have shown great promise as an electrode in glucose sensing devices.²² Because of the

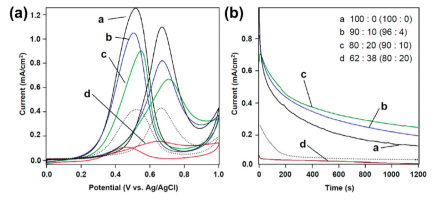


Fig. 12 (a) Cyclic voltammetry and (b) chronoamperometric curves of the Pt-Au mesoporous film at a potential of 0.6 V in the presence of 0.5 M H₂SO₄ containing 0.5 M methanol. The currents were normalized to the ECSAs obtained from CV curves in 0.1 M NaOH solution. The currents were compared to the ECSAs calculated from CV curves in 0.1 M NaOH. The product compositions (Pt: Au) are given in parentheses, as are the Pt^{4+} : Au³⁺ compositional ratios in the precursor solutions. A bulk Pt film (shown by a dotted line) is used as a comparison to the other samples. Electrochemical deposition at a constant voltage (0.04 V vs. Aq/AqCl) was used to make the bulk Pt film. All the samples had a film thickness of less than 100 nanometers. Reproduced with permission. 40 Copyright 2012 American Chemical Society.

synergistic effect of mesoporous structures and Pt-Au binaryalloy surfaces, they have good catalytic activity for glucose oxidation. The amperometric glucose sensors made from mesoporous Pt-Au alloy films have a good anti-poisoning effect, as well as good stability and reproducibility. These films can retain their activity towards the detection of glucose at least for ten cycles. The film was able to detect up to 352 $\mu A~cm^{-2}$ mM^{-1} glucose with a LOD of 6.0 μ M (S/N = 3) (Fig. 13).²² When it comes to industrial manufacturing, however, the costeffectiveness of using precious metals for electrode fabrication is becoming an issue that needs to be addressed. This drawback can be minimized by introducing cheap metals along with the noble metal. For instance, Au and Cu were coupled to form mesoporous Au-Cu alloy sheets, which reduced the manufacturing costs as compared to monometallic Au-based devices. Moreover, this bimetallic alloy boosted both selectivity and sensitivity due to the synergistic electrical response and catalytic benefits from both components.²³ With great selectivity, sensitivity, and good linearity over a broad range of glucose concentrations with a low detection limit, this mesoporous Au-Cu film exhibits outstanding performance at a reasonably strong negative potential, indicating the removal of interferents. 188 Besides these mesoporous bimetallic alloys, ternary alloys, mesoporous Au-Cu-Ni films, also displayed intensified activity in the electrocatalysis of glucose compared to the pure Au film. 72,157 These trimetallic alloys presented a great deal of stability in retaining their activity in glucose oxidation reactions by minimizing the loss of ECSA in respective reactions. 157

4.4. Biosensors

Apart from use in glucose sensing devices, mesoporous metallic alloys also have application in ultrasensitive detection of nucleic acids such as microRNA (miRNA). miRNA, a kind of small non-coding RNA (21-24 nucleotides) that regulates over 30% of human genome, is involved extensively in almost every

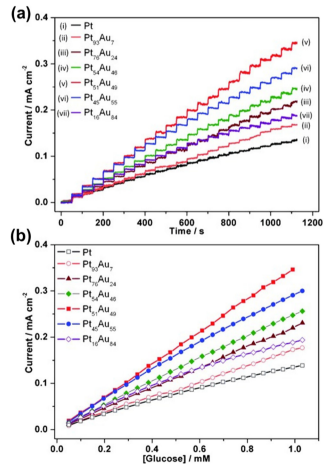


Fig. 13 (a) The dynamic amperometric response of a Pt-Au alloy for the successive addition of various amounts of glucose, ranging from 0 to 1.05 mM at 0.3 V. (b) The corresponding curves of those concentrations. The average values are obtained by using five independent measurements from five different film electrodes. Reproduced with permission.²² Copyright 2013 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim.

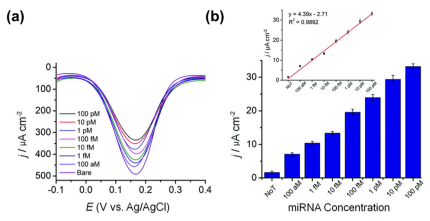


Fig. 14 Assay sensitivity for the Au-Ag bimetallic alloy-based sensor for miRNA. (a) The DPV responses for the Au-Ag film (50:50) sensor for a range of microRNA (miR-9-2) ranging from 100 aM to 100 pM. (b) The relative current is determined by using the bare (non-porous) mesoporous Au-Ag film as a reference. Reproduced with permission. 117 Copyright 2020 Royal Society of Chemistry

cellular process including cell differentiation, genetic expression, and biological development. 189-191 The expression of miRNA changes according to the type and stage of disease, especially cancer, and has been reported to be imbalanced in many disease conditions, either upregulated or downregulated relative to the healthy state. 192 As a result, miRNA expression has been extensively studied as a potential biomarker for the diagnosis and prognosis of a wide range of illnesses, including cancer. 193-195 Mesoporous Au-Ag alloy films have a lot of potential for fabricating portable, high-performance miRNA sensors. 117 These DPV sensors enable ultrasensitive detection of miRNA with an LOD of 100 aM and possess activity in the dynamic range of 100 pM to 100 aM with good reproducibility (RSD, of <5%, for S/N = 3) (Fig. 14). These bimetallic alloys provide several benefits over the current miRNA assays with different cascade signal amplification including high stability, increased adsorption sites for miRNA, enhanced catalytic effect, and therefore, superior sensitivity. 196 Furthermore, the fabrication technique is both cost-effective and reproducible and re-usable. This assay requires a single-step electrochemical detection of isolated target miRNA from the source, requiring less than 1 hour for miRNA detection, making it a good candidate for a portable miRNA sensor for clinical use. It could be used in biological studies, bioanalysis, and clinical diagnosis. Although the potential of the mesoporous nanoarchitecture is utilized for electrochemical biosensing, it could be also applicable for many other types of biosensing applications. For example, the unique properties of the metallic nanostructures might potentially enhance plasmonic phenomena which in turn could be utilized in surface plasmon resonance (SPR) based biosensing platforms.

5. Conclusions and perspectives

In this comprehensive review, we have comprehensively reviewed the current strategies available for the synthesis of mesoporous alloy films. We have discussed the electrochemical

procedures to fabricate various bimetallic and trimetallic alloy films and their corresponding application. To sum up, alloying of noble metals with inexpensive metal species contributes positively to both procedural and functional aspects. Alloying of metals in a bimetallic and trimetallic fashion reduces the overall production cost significantly compared to that of single metal-based catalysts. Alloying also offers highly diversified explorable options to fabricate multi-component alloys with tunable metal constituents. Moreover, the fabrication of binary and ternary alloys by electrochemical methods is highly reliable, cost-effective, and easily reproducible. From functional aspects, electrochemically synthesized multi-metallic mesoporous alloy films exhibit substantial performance boosts in catalytic conversion reactions. On top of that, electrochemically fabricated mesoporous alloy films show high stability, reusability, and durability in participating reactions alongside notable efficiency in resisting catalytic poisoning effects. The high functionality and catalytic performance of electrochemically synthesized mesoporous alloy films (binary and ternary) can be attributed to several factors including highly ordered mesopores across the film, evenly distributed mesopores of the same size throughout the entire cross-sectional area of the alloy film, absence of any cracks and voids, larger surface area provided by constituent metal species, synergistic effects of participating metal species, availability of abundant multifunctional active sites for the guest species, variability in the electronic structure and strain effects, and lastly high stability and durability. Electrochemical methods also offer customizability during the fabrication process to tailor the final film composition and subsequent catalytic performance of the resulting alloys. Adjustment parameters include the compositional ratio of participating metal precursors, size and type of the micellar templates, duration of electrodeposition, etc. Mesoporous metallic alloys fabricated by electrochemical methods are suitable for a wide range of applications including electrocatalytic activity in HER, MOR and electrocatalytic signal amplifications for glucose, protein and nucleic acid sensor. Bi- and tri-metallic mesoporous alloys have been reported to

function as non-enzymatic glucose sensors with high sensitivity, accuracy, and low detection limit compared to the monometallic counterparts and conventional glucose sensors. The versatility and applicability of mesoporous binary and ternary alloy films explored to date are just the tips of the iceberg. Despite the fact that mesoporous metal alloys have been extensively studied in HER and MOR, the application of these films in the avenue of biological sciences such as detection of various types of nucleic acids, biosensing of epigenetic modifications (DNA methylation and RNA methylation), detection of post-translational protein modification, protein purification, biomolecule detection, and so on is still unanswered. We envisage that more focus should be directed towards research exploring alloying of earth-abundant elements and translating their application in biological studies and industrial-scale electrocatalysis.

Author contributions

J. Bashir, B. Chowdhury and R. R. Kathak contributed equally.

Conflicts of interest

There are no conflicts of interest to declare.

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