Review Article

Current Trends in Nanoporous Anodized Alumina Platforms for Biosensing Applications

Ganesan Sriram,1 Pravin Patil,1 Mahesh P. Bhat,1 Raveendra M. Hegde,1 Kanalli V. Ajeya,1 Iranna Udachyan,1 M. B. Bhavya,1 Manasa G. Gatti,1 U. T. Uthappa,1 Gururaj M. Neelgund,2 Ho-Young Jung,3 Tariq Altalhi,4 Madhuprasad,1 and Mahaveer D. Kurkuri1

1Centre for Nano and Material Sciences, Jain University, Jain Global Campus, Bangalore-562 112, Karnataka, India
2Department of Chemistry, Prairie View A & M University, Prairie View, TX 77446, USA
3Department of Environmental Energy Engineering, Chonnam National University, Gwangju 500-757, Republic of Korea
4Department of Chemistry, Faculty of Science, Taif University, Taif, Saudi Arabia

Correspondence should be addressed to Ho-Young Jung; jungho@chonnam.ac.kr, Madhuprasad; madhuprasad@jainuniversity.ac.in, and Mahaveer D. Kurkuri; mahaveer.kurkuri@jainuniversity.ac.in

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Pristine aluminum (Al) has received great deal of attention on fabrication of nanoporous anodized alumina (NAA) with arrays of nanosized uniform pores with controllable pore sizes and lengths by the anodization process. There are many applications of NAA in the field of biosensors due to its numerous key factors such as ease of fabrication, high surface area, chemical stability and detection of biomolecules through bioconjugation of active molecules, its rapidness, and real-time monitoring. Herein, we reviewed the recent trends on the fabrication of NAA for high sensitive biosensor platforms like bare sensors, gold coated sensors, multilayer sensors, and microfluidic device supported sensors for the detection of various biomolecules. In addition, we have discussed the future prospectus about the improvement of NAA based biosensors for the detection of biomolecules.

1. Introduction

In recent years, metal oxide based nanomaterials are emerging in various areas such as thermal barrier [1], gas sensor [2], solar cells [3], drug delivery [4], and biomedical applications [5] due to ease of preparation and possibility in tailoring chemical and physical properties. Nanoporous oxide materials possess tremendous applications in water purification [6], biosensors [7], therapeutics [8], optical sensors [9], oxygen carrier [10], battery technology [11], and bioimaging [12]. Overall, nanoporous anodized alumina (NAA) has several advantages such as high surface area to volume ratio, thermal stability, chemical stability, electrical insulation, formation of unique pore structures, high porosity, self-organized pore structure, being nontoxic and bioactive, and biocompatibility. In addition, systematic arrangement of nanopores is of great interest, especially pores with high aspect ratios between pore diameters to pore length for various applications such as separation, drug delivery, and biosensing. Due to this, geometrical arrangements of pores with high surface area are promising candidates in biosensing for the detection of extremely rare/important biological entities.

Keller et al. have developed closely packed hexagonally arranged duplex structured NAA with pores and barrier type of oxide layer formation by anodic oxide coating through electrolytes [13]. The most popular fabrication method to synthesis NAA is electrochemical anodization, which has wide variety of applications in areas such as filtration process [14], biosensor [15, 16], drug delivery [17], corrosion resistant [18], oxygen sensor [19], catalysis [20], photocatalytic [21], cancer therapy [22], refractometric [23], interferometric [24], DNA sensor [25], bone implant [26], separation [27], and fluorescence detection [28]. The growth of porous structure depends on the ionic transport in anodic oxides under the
influence of electric field by anodic polarization in a suitable electrolyte. The pore geometry (pore shape and diameter) depends on the processing parameters such as electrolyte solution, pH, anodization voltage, current density, time of anodization, and temperature. However, precise pore diameter formation in the anodization process of pristine aluminum (Al) under applied electric field is completely not understood yet.

Al can be electrochemically anodized by two-step anodization process to get nanopores [29, 30]. In anodization process, its surface is electrochemically oxidized in an electrolyte solution which leads to the formation of an oxide layer. The anodization can be done by applying voltage between an Al electrode (anode) and Pb and/or Pt electrode (cathode) arranged parallel to each other. The electric field is equally dispersed on its surface where pores are created on the surface of NAA [31]. The highly ordered porous structures are fabricated under low temperatures with strong acid electrolyte solutions (oxalic acid, phosphoric acid, and sulfuric acid) which is called the mild anodization process. Further, these self-ordered pore arrangements with narrow size distribution and high aspect ratio can be developed by applying different potentials, for example, at 25 V with sulfuric acid [32], 40 V with oxalic acid [30], and 195 V with phosphoric acid [33] leading to the formation of pores of diameters 65, 100, and 400 nm, respectively. Despite being advantageous to form precise pores, the main disadvantage of mild anodization or self-ordered arrangement is that the rate of oxide growth is slower due to low current density and high processing time. On the other hand, hard anodization process can overcome the drawbacks of mild anodization process where higher current densities and faster processing steps are involved and the whole process is mechanically robust [34]. Recently Losic and Santos reported the fabrication of NAA by soft and hard anodization techniques for structural changes of pores and optical properties for specific sensing applications [35]. However, by using acid solutions porous structures can be created in NAA; on the other hand, by using neutral solutions barrier (nonporous) type of NAA is created [36]. Further, Lee and Park published a review on the formation of structural changes of porous and nonporous (barrier) types of NAA by anodization (Figure 1) [37].

In addition to that, NAA has been used in separation studies such as protein [38], molecular release [39], and transport [27]. This process offers self-organized growth of nanopores from top to bottom which provides highly ordered NAA structure. The fabrication of nanopores in NAA of different structures such as funnel [40], multilayered [41], hierarchical [42], and three-dimensional [43] has also been reported. Recently, Santos et al. developed NAA with inverted funnel structure; this type of multistratified structures of cylindrical nanopores varies in pore diameter from top to bottom [44]. The NAA can also be fabricated in wide variety of nanostructures such as nanotubes, nanofibers [45], and nanodots [46].

Owing to the abovementioned importance and wide range of NAA applications herein, we are reviewing fabrication and applications of NAA in biosensing arena. In this article, the most recent and current research on NAA and gold coated NAA (Au-NAA) with their key role in biosensing applications have been covered. In particular, our focus is on fabrication of NAA by anodization process and their specific applications in biosensing with detailed description on fabrication conditions, working principles, and mechanism of biosensing. Finally, the future prospectus of NAA in the area of biosensing has been discussed.

2. Fabrication of Nanoporous Anodized Alumina (NAA)

In general, NAA can be developed by two-step anodization method (Figure 2) [47, 48]. As discussed in Introduction, by varying parameters like applied current, concentration of acid solutions, anodization time, and temperature, one can tailor the NAA properties. For example, Yin et al. used same method [63]; at first the Al film was electrochemically polished in a mixture of ethanol (EtOH) and HClO$_4$. In the next step, Al was anodized with 0.3 M oxalic acid at 10° C and at 40 V for few hours to obtain nanoporous structures of NAA. Further, second anodization was carried out in the same way to remove the barrier layer which is created at the bottom of nanopores by using 0.5 M phosphoric acid or by using dry etchants like chlorine-based gases. In this method highly ordered nanopore arrays in thin alumina film on silicon substrate were obtained.

Similarly, Lo and Budiman fabricated NAA by using impure Al foils which can be used in large scale applications whereas the highly pure foils have limited applications in nanoelectrophotonics. The impure Al foil was taken as anode and copper as cathode [64]; further, 5 × 5 mm pieces of impure Al foil were mounted on the glass slides and these were then anodized in 0.3 M oxalic acid for about 18 h at constant voltage of 40 V. The oxidized layer was etched using 0.5 M phosphoric acid at 80° C for 1 h. The same material was anodized by using above conditions for 15 min to develop nanopores throughout the surface. Thus, obtained samples were removed and floated over saturated HgCl$_2$ and further treated with H$_3$PO$_4$ to remove the deposited barriers to get NAA. To compare, ultrapure Al foils were used and it was found that all the properties such as growth rate and chemical
composition were similar with that of NAA produced from impure Al foils except for the pore ordering (morphology). However, the pore consistency and order can be controlled by electropolishing the impure Al foil.

Nagaura et al. proposed a multistep anodization and leaching process to get three dimensional structured films [65], in which Al foils were used as working electrode in anodization process and carbon electrode as counter electrode. Each anodization step was followed by dipping the substrate into phosphoric acid for pore widening, which was called leaching process. The anodization was conducted in phosphoric acid solution at 10°C with stirring and varying current to get multistep structures which is followed by leaching process after every anodization step with different time interval of dipping.

Similarly, Hussain et al. fabricated uniform NAA to fabricate tungsten trioxide (WO$_3$) nanowires [66]. In this process, Al foils were cleaned with acetone to remove the dirt and greases which were further subjected to annealing at 500°C for 5 h at heating rate of 10°C/min. Further, the oxides were removed by dilute oxalic acid and rinsed with deionized water. In next step, electrically polished foils were kept in a mixture of EtOH and perchloric acid at a voltage of 12 V for 3 min at 100°C. Further, two-electrode cell system was used to fabricate NAA using graphite as counter electrode. Anodization was carried out at 40 V for 120 min at 50°C in 0.3 M H$_2$C$_2$O$_4$ electrolyte. The oxide layer of NAA substrate was removed by washing with H$_2$Cr$_2$O$_7$ and H$_3$PO$_4$ for 30 min. Further, it was again anodized at a voltage of 40 V for 120 min at 50°C in 0.3 M H$_2$C$_2$O$_4$ acid. After this process, NAA was immersed in H$_3$PO$_4$ for 90 min to increase the size of pore width. The fabricated NAA has been used as a template to fabricate WO$_3$ nanowires by electroless deposition method. NAA was sonicated under the mixture of PdCl$_2$ and HCl used in the wt% of 0.1 and 1, respectively, for 3 min which is followed by dipping the template in deionized water. The deposition bath was used to form nanowires by NAA template in which sodium tungstate, hydrazine hydrate, sodium potassium tartrate, and acetic acid were used as a composition. With this, WO$_3$ nanowires were formed in the pores of NAA.

For the formation of unidirectional NAA structures Na et al. [67] used pure 1050 Al sheets after pretreatment to clean the surface before the anodization process. Al samples were anodized in dilute H$_2$SO$_4$ electrolyte solution using Pt cathode at a current density of 3 A/dm$^2$. In the next step, anodization was carried out at different electrolyte temperatures such as 2, 10, and 15°C for 5 to 6 h. After this process, acid resistant tape was removed by acetone washing to form porous structures on surface. The thickness and average diameter of unidirectional nanopore were found to be 230–240 μm and 6–24 nm, respectively.

In another work, the synthesis of carbon nanotubes in NAA film by H$_2$SO$_4$ or H$_2$C$_2$O$_4$ acid was achieved by Sui et al. [45]. Polished Al plate was anodized at different anodic voltages in 0.16 M H$_2$C$_2$O$_4$ or 1.2 M H$_2$SO$_4$ solution for 25 min by vigorous stirring at 15°C to get porous NAA template, which was then cleaned and kept for boiling (15 min). Pairs of alumina templates with or without boiling in water were put into tubular stove with gas flow mixture of N$_2$C$_2$H$_2$ at the rate of 100 mL/min at a particular temperature for 2 h. Further, Al was dissolved in 10% NaOH solution to release nanotubes and nanofibers. Immediately, these were thoroughly washed and dispersed in pure EtOH solution by sonication method to obtain pure nanopore structures in the form of carbon nanotubes on NAA. Furthermore, Tsuchiya et al. developed self-organized NAA in neutral electrolytes [68]. Al foils (0.5 mm) were degreased by sonicating in acetone, isopropanol, and methanol followed by rinsing in deionized water and dried under nitrogen atmosphere. Anodization was done in

![Figure 2: (a) Experimental set-up for fabrication of NAA by two-step anodization process and (b) the geometric structure of NAA (d$_{int}$: interpore distance, d$_{p}$: pore diameter, and L$_{p}$: pore length) [47, 48].](image)
a conventional cell with three electrodes; Pt as counter electrode and Haber-Luggin capillary electrode with Ag/AgCl were as reference electrode. The samples were assembled in electrochemical cell by pressing against an O-ring. The electrolytes such as 1 M (NH₄)₂SO₄ solution with fluoride ions and deionized water were used in this experiment. Anodization was done at high voltage potential which consists of potential from open circuit potential to certain fixed potential. Further, the samples were rinsed in deionized water and dried in nitrogen stream. The pore diameter of NAA was obtained at different potentials in acid solutions ranging from 65 to 500 nm.

Fabrication of NAA with ideally ordered channel arrangements was reported by Asoh et al. [69]. In this, the electropolished master mold was placed on pure Al sheet below 10°C for about 4 min in a mixed solution of HClO₄ and C₂H₇OH. An array of concaves on Al surface was formed. Further, anodization of Al was done by 0.3 M H₂SO₄ and/or H₂C₂O₄ under constant voltage. Later, anodic oxide film was removed from Al substrate in saturated HgCl₂ solution. The selective removal of barrier layer (down part of oxide layer) and pore widening were done by etching the NAA membrane in 5 wt% H₃PO₄ at 30°C. This treatment facilitates better observation of the holes at bottom oxide layer, where uniform diameter of channels (40 nm) and depth of the nanopore (1 μm) were formed.

On the other hand, Zhang et al. fabricated NAA membrane using low potential anodizing process [70]. Highly pure Al foils were cut into small pieces and annealed in a furnace in the presence of air at 600°C. Mixture of acids (H₃PO₄ : HNO₃ : H₂O = 8 : 1 : 1) was taken and Al plates were polished for 30 s at 95°C followed by electrochemical polishing in perchloric acid and EtOH (1 : 4 ratio). Anodization was done in H₂SO₄ aqueous solution under constant potential of 18 V. The Al plate was fixed in acid solution which was separated from cathode of mechanically polished copper plate. All experiments were conducted in 3°C cold water using ice bath. The oxide layer was produced at first anodization step and was removed by submerging the Al plate in chromic acid solution. Second anodization was conducted under same conditions by shortening the duration to 3 h. The NAA were obtained with pore diameters ranging from 6.8 to 19.3 nm at potential of 6 to 18 V.

Wang et al. fabricated NAA nanotubes using pulse anodization; the NAA nanotubes were fabricated by modified anodization process, that is, pulse anodization [71]. Al plate was sonicated in EtOH, ultrapure water, and then electropolished in a mixture of EtOH and HClO₄ 4 : 1 (v : v) at 20°C and 5°C for 3 min. The first anodization was done in an electrolyte solution of 0.3 M H₂SO₄ at 25 V and 6°C. This consists of mild anodization and hard anodization pulses in 0.3 M H₂SO₄ electrolyte and EtOH of concentration ranging from 0 to 25%. Wet chemical etchants were used to remove remaining Al substrates in 0.2 M CuCl₂ and 6.1 M HCl solution. Through pulse anodization, the pore length and diameters of NAA were found to be 500 and 90 nm, respectively.

Friedman et al. prepared series of highly porous and pure NAA [72]. A flat cell was used with a Pt mesh counter electrode, and the Al was exposed to electrolyte at 0.25 ηcm². At first anodization step, the NAA was anodized for 2 h and kept in 5% H₃PO₄/5% chromic acid solution for overnight which is followed by etching to provide a barrier layer, thus resulting in the formation of small porous structures on the surface of Al. Further, it was reanodized at the same condition as in the first step but with shorter time interval. So, the above methods emphasize the fabrication of different types of NAA with different pore diameters and pore lengths for the detection of immobilized biomolecules on their surface. In addition, these key parameters enhance the optical properties and rapid sensitivity in the detection process of biomolecules. Consequently, the structural parameter and fabrication conditions of the NAA become crucial platform for biosensing application.

3. Applications of NAA in Biosensing

Recently, NAA based materials have attracted considerable attention in biosensing due to possibilities like immobilization of target molecules on nanopores, optical sensing enhancement by coating Au layer on NAA surfaces, low concentration detection, biocompatibility with living cells due to Au coating, portability, and so forth. The recent reviews on NAA based biosensing date back to 2014 [73–76]; thus, there is a huge scope for the current review article which covers latest research works and current trends in NAA based biosensing applications. The following applications are categorized based on the types of transducer for the detection of biomolecules.

4. NAA Biosensor

4.1. Detection of Glucose. The barcode system enhances the optical biosensing effect of NAA for oxazine dye and glucose using photoluminescence (PL) spectroscopy [77]. Enhancement in the PL barcode system for optical biosensor was observed for NAA structure with wide pore diameter (30–71 nm) and controlled pore length (5–12.4 μm). The barcodes were produced with respect to different wavelength of PL spectra. The wide pore diameter and controlled pore lengths were obtained by wet chemical etching and various anodization processing times. It was found that PL barcode system depends on the pore diameter and lengths. In a control experiment, no PL spectrum was observed for the pore length larger than 25 μm with 5 h anodization time. In this study pore lengths, such as 5, 8, and 12.4 μm, resulted in better PL intensities. Intensity of PL spectra was increased with wide pore diameter, due to decrease in surface thickness of substrate. However, PL intensity has been decreased while increasing the etching time for longer pore diameters. Further enhancement in PL intensity for larger pore diameter was exhibiting better sensing effect for oxazine dyes and glucose biosensors.

Furthermore, Santos et al. studied the effect of pore length and diameter of NAA for biosensing applications [78]. The study showed effect of pore diameter and pore length with respect to particular reflection in NAA. Samples
with different pore diameters with same pore lengths were considered with same geometrical characteristics. The wider pore showed greater reflectivity due to bigger pore diameter and larger pore gives lower reflectivity due to large pore length. The reflectivity studies have been used to predict the biosensing mechanism of NAA. Furthermore, Santos et al. reported the comparison of biosensor material for the detection of D-glucose and L-cysteine using PL and reflectometric interference spectroscopy (RIFS) [79]. NAA substrate was fabricated by two-step anodization process with pore length and pore diameters of 1.8 to 12.7 µm and 29 to 74 nm, respectively. The detection of D-glucose and L-cysteine under nonspecific and specific adsorption conditions by NAA was carried out using PL and RIFS systems. The better sensitivity for both the biomolecules was observed for NAA by PL due to their photoactivity compared to RIFS. Sensing of glucose and L-cysteine was achieved at low concentration by NAA, that is, 0.01 M and 0.005 M, respectively, using PL.

In another work, Kumeria et al. developed NAA rugate filters (NAA-RFs) for real-time detection of D-glucose by reflection spectroscopy with flow cell [15]. Reflection spectroscopy is one of the optical instruments with various benefits such as small size, being portable, and low cost, compared to other optical instruments. Four different types of NAA rugates, that is, NAA-RF 1, NAA-RF 2, NAA-RF 3, and NAA-RF 4, were fabricated. The detection of D-glucose in flow cell was observed by change in refractive index (RI) of NAA rugate filter with respect to adsorption of D-glucose into nanopores. The RI was measured initially by using distilled water which was sent into flow cell followed by D-glucose solution. Further, this D-glucose was immobilized on NAA-RF 1 where RI has been increased for D-glucose and decreased for distilled water. Similar procedure was applied for other three NAA rugate filters. However, better sensitivity was found for NAA-RF 2 compared to other three NAA-RFs materials for D-glucose. The schematic diagram explains the optical system for detection of D-glucose by combining NAA-RFs and reflection spectroscopy and sensing plot for different concentration of D-glucose with respect to change in effective RI of NAA-RFs using reflection spectroscopy (Figure 3).

4.2 Detection of Binding of Antibody and Antigen. Takmakov et al. fabricated NAA for the detection of streptavidin through covalently bonded biotin on sensor surface using fluorescence microscopy [80]. By using aminosilane-succinimide chemistry [81], the biotin molecules were immobilized covalently on NAA surface. Fluorescence microscopy was used to detect streptavidin molecules on biotin bonded sensor surface. The fabrication process and sensing mechanism are explained as follows: Al film was coated on glass substrate through mask by ion beam coater with film thickness of around 600 nm. The anodized Al film showed pore diameter and thickness of 20 nm and 60 µm, respectively. The biotin molecules were immobilized on sensor surface by NAA which were rinsed in 3-aminopropyltrithoxysilane (APTES) and later washed with acetone followed by drying in oven. Biotin layer was formed on sensor surface by mixing with dimethyl sulfoxide (DMSO) solution of biotin N-hydroxysuccinimide ester and acetone. The covalently
bonded biotins were analyzed on surface by FTIR. Further, biotinylated substrate was exposed to streptavidin modified Alexa Fluor 488 dye solution for 5 min. The unbound streptavidin on biotinylated substrate was removed by further washing with phosphate-buffered saline (PBS) solutions. The streptavidin bounded biotin NAA substrate was immersed into water and then analyzed by fluorescence microscope to confirm the binding of biomolecules. The streptavidin molecules were exposed to both materials in which the fluorescence contrast was high for NAA compared to Al film due to high diffusion of streptavidin into porous structured NAA. The sensing of streptavidin was carried out by using fluorescence microscopy where significant increase in fluorescent intensity was observed for NAA (black line) due to covalent bonding of streptavidin-biotin on sensor surface and moreover biomolecular diffusion process was higher in porous surface. In contrast, Al film exhibited very low intensity peaks due to improper binding of biomolecules and was having less diffusion due to nonporous structure of bare metal film.

Fu et al. developed NAA based biosensor for the detection of antigen-antibody detection using SERS method [82]. NAA was developed by the process of Al thin film coated on glass substrate with thickness of 380 nm by thermal evaporation technique and followed by anodization process. Further, silver (Ag) nanoparticles were coated on the surface of NAA to enhance the RI of medium. The immobilized antigen-antibody was obtained on modified NAA substrate by washing with APTES solution followed by GTA solution for the formation of amino and aldehyde groups, respectively. The detection of antigen-antibody recognition on NAA was observed by wave guide assisted SERS system. The obtained human IgG (antigen) was covalently bound with NAA surface followed by FITC-anti-IgG (antibody) which were allowed to immobilize with human IgG modified NAA surface. The immobilization of molecules was analyzed by change in angle dependent reflectivity on NAA surface. The angle dependent reflectivity studies were obtained by processing bare NAA substrate at angle 44.74°, human IgG modified NAA surface has angle of 45.04°, and at the end of FITC-anti IgG modified NAA the angle was 44.42°. The above results show that the angle dependent reflectivity increases with respect to excess binding of molecules on NAA surface. The detection of binding molecules on NAA surface was analyzed by enhanced SERS signals through Ag nanoparticles accumulation.

Kant et al. developed biosensing platform by utilizing the pore dimension of NAA to detect the immobilization of streptavidin (antibody) with biotin (antigen) by electrochemical impedance spectroscopy [49]. NAA has been fabricated with 25–65 nm and 4–18 μm pore size and lengths, respectively. In addition, nanopore arrays were patterned on back side of substrate by photolithographic process. The streptavidin biomolecules were bound on nanopore array structure of NAA surface by soaking it in mixture of streptavidin with PBS solution. The different concentration solutions of biotin molecules were allowed to bind on streptavidin functionalized substrate. However, 0.2 μM concentration of biotin molecules bound with streptavidin showed better sensing performance due to decrease in pore resistance on substrates and increase in pore diameter from 25 to 65 nm. It has been found that sensitivity has not only depended on pore dimensions where size of biomolecules may also influence the enhancement of sensing performance. The size of immobilized biomolecule of streptavidin was about 4–5 nm and the size increased to 6–8 nm due to binding with biotin molecule which resulted in better performance. The electrochemical impedance spectroscopy was performed by four electrode systems to characterize the NAA biosensor. The electrochemically impedance biosensor performed by change in surface charge of NAA binding to both biomolecule and pore resistance of substrate. Two gold foils as working and counter electrodes with Ag/AgCl were used as a reference electrode. The NAA substrate was sandwiched between two cells and contacted to reference electrode from each side of cells. The working and counter electrodes were placed at the end of two cells. An equivalent circuit was used to balance the resistance and capacitance of the NAA to analyze the streptavidin bonded biotin on NAA. The sensitivity of NAA has been enhanced by decreasing pore resistance with respect to change in conductivity of sensor surface. The larger pores with longer pore length are not suitable for the detection of biomolecules. Thus, the pore resistance increased when the pore length was increased from 4.5 to 18 μm. The pore resistance depends on the pore length and it was found that no response/sensitivity was observed for the pore length above 10 μM. The detection limit was observed using different solutions of biotin in the range of 0.2 to 10 μM by binding with streptavidin on NAA substrate, but in the concentration of 0.2 μM it showed better sensitivity for the binding of biotin and streptavidin molecules on NAA substrates. The fabrication and formation of nanopore arrays on NAA surface by lithographic technique and graphs are shown in Figure 4. The graph represents the pore resistance dependent on different concentration of biotin solution with respect to different pore diameters and pore lengths obtained by NAA biosensor.

4.3. Detection of Protein. Alvarez et al. reported real-time detection of binding protein-A with rabbit, anti-sheep, and sheep immunoglobulin on the surface of NAA with respect to change in optical thickness using fluorescence microscopy [50]. The fabricated label-free NAA biosensor has structural dimensions of 60 nm and 6 μm in diameter and thickness, respectively. The sensor surface modification by protein-A has been tested for interaction with rabbit, sheep, rabbit, anti-sheep and chicken's immunoglobulin. The binding interactions were carried out in PBS solution with 0.1 mg/mL biomolecules concentration for the biosensing applications. Schematic diagram in Figure 5 represents the protein-A modified sensor surface and step by step binding mechanism of rabbit anti-sheep and sheep's immunoglobulin biomolecules with protein-A. The sensitivity has been measured in terms of time versus change in optical thickness (OT) of sensor surface modified protein-A with rabbit, rabbit anti-sheep, sheep, and chicken immunoglobulin. Optical thickness has been rapidly increased with respect to time on NAA sensor surface for protein-A interacting with rabbit anti-sheep and
sheep immunoglobulin biomolecules. However, no binding affinity was observed for rabbit and chicken immunoglobulin biomolecules (dark base lines) due to weak interaction with sensor surface.

In addition to that, Lau et al. fabricated NAA for the detection of bovine serum albumin (BSA) through pore widening using optical waveguide spectroscopy [83]. The immobilized BSA on NAA surface was observed for reflectivity to confirm the detection. High sensing performance was attained through the pore widening on NAA surface. The reflectivity increased with respect to increase in the sensitivity of material. The enhancement in sensing performance...
was obtained by processing film with uniform porous structure.

4.4. Detection of DNA. Pan and Rothberg developed immobilization of single stranded DNA (ssDNA) oligonucleotides on NAA and the fabricated materials were functionalized as a reflective biosensor to detect complementary DNA at area of 1 nmol/cm² [84]. The fabricated structures of NAA were of 6 µm and 40 nm in thickness and diameter, respectively. The fabricated sensor material has very high surface area which could easily accommodate DNA oligonucleotides on NAA surfaces. The detection of biomolecules was based on change in reflectivity on the surface of sensor material.

Takmakov et al. studied the ionic conductance via shrunk porous alumina and ssDNA biosensing by using porous material [85]. The NAA substrate was fabricated with the structural dimensions of 60 nm and 10 µm pore diameter and depths, respectively. Further, NAA pore diameters were shrunk by hydrothermal treatment. The process involves three steps, formation of shrunk porous structures on sensor material by anodization, the reduction in the size of porous structures by hydrothermal treatment, and the detection of DNA oligomers by porous materials. The ssDNA biomolecules were immobilized on sensor surface by aminosilane-glutaraldehyde chemistry [86]. The immobilization of ssDNA inside the sensor material decreases the pore resistance due to increase in the surface charge density of sensor material with negatively charged ssDNA. The immobilized ssDNA was hybridized with double stranded DNA (dsDNA) to increase the pore resistance. The dsDNA was denatured using urea which liberates complementary DNA strands to recover the initial pore resistance of immobilized ssDNA. The DNA detection has been achieved by the conduction variation with or without hybridization of DNA in internal nanopores of sensor material through electrical impedance studies.

Hotta et al. developed the multilayer structure polyelectrolytes coated NAA surface for the detection of DNA (single stranded probe DNA) and hybridized DNA using infrared reflection adsorption spectroscopy (IRRAS) [51]. The Al thin film was deposited on glass substrate by thermal evaporation followed by anodization to get pore diameter of 40 nm. Further, the polyelectrolyte solutions (poly (acrylic acid) and poly (allylamine hydrochloride)) were coated on NAA substrate using layer by layer coating process to form a protective layer during buffer solution immersion process.

Figure 5: (a) The schematic diagram of binding affinity of biomolecules on NAA surface and (b) the time dependent change in optical thickness (OT) graphs showing the binding interaction of protein-A, rabbit, rabbit anti-sheep, sheep, and chicken immunoglobulin [50].
Detection of DNA-DNA hybridization on NAA substrate by optical wave guide spectroscopy was carried out as follows: initially streptavidin was immobilized with biotin on NAA surface to detect DNA which showed increased red shift upon binding. Further, this immobilized DNA on NAA surface was bonded with noncomplementary (T2) target DNA. Thus, red shift has been increased due to complementary target DNA (T1) added with probe due to hybridization. Better sensing detection was found from complementary target DNA (T1) probe on NAA surface due to DNA hybridization which was observed using optical wave guide spectroscopy. Schematic representation of polyelectrolytes-NAA film and DNA-DNA hybridization assay using polyelectrolytes-NAA is shown in Figure 6.

4.5. Detection of Viral Particle. Moon et al. developed NAA to capture and align procapsid (phi29 viral) particles using fluorescence microscopy [87]. Al foil was anodized to prepare the diameter and interpore distances of 75 and 105 nm, respectively. The pore structure was modified by coating of Al2O3 film using atomic layer deposition (ALD) with average pore diameters of 38 nm. The surface of NAA was cleaned using 3 (trimethoxysilyl) propyl aldehyde (TMSPA/TMS) solution. Later, substrate was incubated in phi29 viral and followed by rinsing in TMS buffer solution to remove unwanted phi29 viral particles on surfaces. Later, DNA package on NAA substrate was tethered to capture phi29 viral particles through polishing and centrifugation. By polishing, the chemically modified NAA substrate was incubated in procapsid solution so that high numbers of phi29 viruses were bound into nanopores and small amounts of viruses were out of the nanopores. By centrifugation, the pore sizes were shrunk to 38 nm by ALD method: actual size of phi29 viral particles was 45 nm. The attractive capture behavior was observed via centrifugation for phi29 viral particles on DNA packaged NAA substrate. The nanopore sizes were less than viral particle sizes; thus, it was easy to be captured onto substrate. The fluorescence microscopy was used to detect the viral particles on NAA substrates.

4.6. Detection of Vitamin C. Chen et al. developed the photonic structure of biomimetic based NAA with distributed Bragg reflector for the sensing of vitamin C molecules [52]. Different anodization process was used to fabricate NAA with distributed Bragg reflectors by galvano-static pulse anodization process. However, excellent sensing performance of vitamin C was observed for NAA prepared with anodization period of 675 s at 3 °C with current ratio of 6:1 compared to other anodization parameters. Schematic representation of vitamin C molecule captured by APTES and GTA solution on NAA distributed Bragg reflectors and sensing results of vitamin C molecule with respect to change in effective optical thickness (OT) are presented in Figure 7. Surface functionalization of sensor involves the APTES modified NAA distributed Bragg reflectors with internal surface hydroxylated by using H2O2 solution, further modified with APTES using Chemical Vapor Deposition (CVD) process, and later treated with GTA solution to activate the amine groups of APTES. Thus, modified NAA distributed Bragg reflectors were used to adopt vitamin C molecules on internal surfaces. The RIS has been used to detect different solutions of vitamin C molecules on sensor surfaces with respect to change in effective optical thickness of sensor substrate.

4.7. Detection of Enzyme. NAA has been developed for the detection of glucose oxidase (GOx) using UV-vis spectroscopy by Darder et al. [88]. The pore diameter and thickness of NAA pores were in the range of 30–80 nm and 33–150 μm, respectively. The GOx was immobilized on NAA surface and was fixed between two disks; the acetate
buffer solution was filtered through NAA substrate which was placed in between the disks, followed by the GOx with acetate buffer and chitosan solutions. The lack of chitosan covering on GOx modified NAA surface may decrease the enzyme activity due to leakage of glucose oxidase from the surface. Outer protecting layer of chitosan can permeate the water molecules and allows ions to transport and bind to protein. The UV-vis spectroscopy has been used to analyze the activity of immobilized GOx. The solutions of 5 mM potassium ferrocyanide and 40 mM glucose were mixed in 0.1 M Tris/HCl buffer for analysis. Later, 40 μM GOx was added to the prior solution for enzymatic reaction. The procedure was repeated to attain the linear relationship between the absorbance and GOx reactions. The platinum electrode holds the GOx modified NAA surface for glucose determination. The increase in pore size leads to increase in the enzymatic activity and shows good response even at low concentration. The response was very less for higher concentration. The response was very less for higher concentration.

Recently, polyelectrolytes based layer by layer coating on NAA for the biosensor applications was explored. Krismastuti et al. fabricated layer by layer coating of polyelectrolytes on NAA for the bacterial enzyme detection of proteinase-K using interferometric reflectance spectroscopy (IRS) [53]. Three different thicknesses of NAA such as 1.3, 3.0, and 4.4 μm with pore diameter, interpore distance, and wall thickness of 30 ± 1.7, 100 ± 1.5, and 70 ± 1.8 nm, respectively, were fabricated. The aim was to increase the thickness of porous layer to influence the easy detection of the enzyme. Polyelectrolytes like poly (sodium-4-styrenesulfonate) (PSS) and poly-L-lysine (PLL) have been used as negatively and positively charged layers, respectively, for layer by layer coating onto the sensor surface. Instead of gold coating on alumina surface polyelectrolytes were coated due to several advantages such as ease of availability and low cost and can play a vital role in increasing the RI on sensor surface. The fabrication process of layer by layer deposition of polyelectrolytes on to the sensor surface is as follows: dust free NAA was silanized by APTES, followed by 7–10 nm thick gold film onto NAA substrate to prevent the pore blocking. The second step is layer by layer coating of polyelectrolytes using custom designed flow cell and coating was monitored by IRS. Separately PSS and PLL were taken in NaCl solution: PSS solution, followed by PLL solution at a flow rate of 0.2 mL/min, was injected alternatively into the sensor surface. Thus, multilayer coating of opposite charges of polyelectrolytes was achieved on the NAA surface. The polyelectrolytes coated NAA sensors of different thickness were used to detect bacterial enzyme of proteinase-K by IRS spectroscopy system which is shown in Figure 8. The enzyme exposure to substrate of thickness 1.3 μm shows the effective optical thickness (EOT) of 1.9 nm and after washing sample with Mili-Q water the EOT was blue shifted to 7.1 nm. Proteinase-K enzyme was also exposed to substrate of 3.0 μm thick which reveals that EOT was blue shifted to 1.8 nm and after washing with water it was shifted to 1.9 nm. However, proteinase-K exposure on the substrate of 4.4 μm thick showed red shift in EOT which is due to polyelectrolytes being diffused deep inside the nanopores to form high thick sensor so that enzymes were occupied easily into porous layers and hence increasing the RI of sensor substrate, compared to those of 1.3 and 3.0 μm thick sensor.

A simple and smart detection by using trypsin immobilized NAA through PL system was developed by Santos et al. [54]. NAA was fabricated by two-step anodization method; the average pore size and lengths were of 40 nm and 7.1 μm, respectively. The size of trypsin biomolecule was 3.8 nm and hence molecules were easily accommodated and immobilized.
on the nanopores. The process of trypsin immobilization in nanopores was made possible by silanization process: APTES and glutaraldehyde (GTA) were mixed together with NAA, and later trypsin biomolecules were allowed to bind via amino groups on porous substrate by the incubation process. The schematic diagram of different stages of immobilized trypsin molecules on NAA and sensitivity plot of trypsin by change in effective optical thickness with respect to different stages on NAA surface by PL spectroscopy is shown in Figure 9. The detection of immobilized trypsin has been done by observing change in effective optical thickness on sensor substrate by PL. Effective optical thicknesses of bare NAA, APTES coated NAA, GTA-modified NAA, and immobilized trypsin on NAA were found to be 0.1, 0.6, 1.2, and 1.8%, respectively. From the above observation, it has been confirmed that there was an obvious increment in effective optical thickness of NAA.

Invertase is one type of enzyme which is used for the production of sugars and used for hydrolysis process like inverting sucrose into glucose and fructose. Thus, NAA biosensor was fabricated and applied to real-time detection of invertase enzyme using Surface Plasmon Resonance (SPR) sensor by Dhathathreyan [55]. For the enzyme immobilization, the substrate of pore diameter and interpore distances of 60 and 100 nm, respectively, were used. The uncontaminated NAA substrate was kept with invertase enzyme solution.
for 1 h at required pH value. The enzyme immobilization activity was tested in the experimental flow cell by gold coated glass plate and SF10 prism set-up sealed with Teflon cell to impound tested solution. The immobilized invertase and its interaction with sucrose were analyzed by calculating the change in RI on substrate using SPR signal as shown in Figure 10.

On the other hand, gelatin-modified NAA was fabricated for the real-time detection of trypsin using RIfS by Nemati et al. [56]. Gelatin molecules play vital role for chemical sensing and better selectivity towards trypsin molecules. The NAA substrate was gold coated with 5 nm thickness using sputter deposition, and the APTES and GTA solutions were used to modify the surface of NAA for the formation of amino groups. After this, gelatin molecules were allowed to immobilize on inner surface of the pores. The different concentrations of trypsin enzyme (0.125, 0.3, 0.5, 0.6, and 1 mg mL\(^{-1}\)) were used to analyze the sensing performance by using RIfS system. The effective optical thickness was changed with respect to different concentration of trypsin molecules. However, higher concentration of trypsin molecules (1 mg mL\(^{-1}\)) showed fast response in RIfS. More concentration of gelatin molecules immobilized on the pores provides faster response to trypsin molecules. Hemoglobin has been used instead of gelatin as a chemical sensing element that also shows faster sensing performance. Schematic diagram of enzyme detection by gelatin-modified NAA using RIfS set-up and also different concentration of analyte solution leads to change in effective optical thickness in real-time which is shown in Figure 11.

5. Gold Coated NAA (Au-NAA) Biosensor

5.1. Detection of Protein. While integrating NAA into the SPR system, the RI at the close proximity of the sensing platform plays vital role. Thus, Hierro-Rodriguez et al. have extracted the RI values of different NAA templates which are useful in developing SPR and long range SPR sensors [89]. Similarly, Kim et al. developed an Au-NAA biosensor for the label-free detection of aptamer-protein binding interaction on sensor surface using combination of localized surface plasmon resonance (LSPR) and interferometry [90]. The obtained uniform porous structured NAA substrate had pore diameters and
interpore distances of 60 and 120 nm, respectively. The size of pore diameter was reduced to 47 nm by coating of Au thin film on NAA surface by thermal evaporation technique with thickness of 15 nm for the enhancement in optical properties on their surface. The aptamer-I was immobilized on NAA surface by rinsing NAA with thiolated aptamer-I and 6-mercaptohexanol solutions. Further, aptamer-I modified NAA substrate was rinsed with mixture of thrombin and PBS solution. After processing, aptamer-II solution was interacted with aptamer-I-thrombin modified NAA surface. The RI was high for aptamer-I with NAA compared to bare Au-NAA. The obtained high concentration of aptamer-I was immobilized on Au-NAA which was red shifted and was observed through interferometry system. Later, aptamer-I-thrombin immobilized Au-NAA surface revealed that the red shift was due to enhancement in RI of NAA. However, nonspecific binding of aptamer-I and aptamer-II on Au-NAA surface was observed whereas aptamer-I-thrombin on Au-NAA surface resulted in better binding interaction. The RI depends on the binding interaction of biomolecules on sensor surfaces. The saturation state was attained for the concentration of thrombin more than 100 μM; above this concentration no response was found from interferometry and LSPR signals.

Figure 11: (a) Sensing of enzymes by gelatin-modified NAA using RIfS system, (b) surface chemistry used in enzymatic sensor where gelatin acts as the chemical sensing molecule for the enzyme, (c) the effective optical thickness changes in gelatin modified NAA with respect to the intensity using RIfS, and (d) real-time sensing of enzyme [56].
In recent years, there have been increased interest on developing label-free optical biosensors based NAA. These types of label-free optical biosensors are exhibiting high throughput in medicinal applications. The Au-NAA bilayer substrate as label-free optical biosensor for BSA has been developed by Macias et al. [16]. The bilayer structure of NAA has been prepared with different porosity with pore diameter and depths of 68 nm and 1.9 μm as well as bottom layers with 33 nm and 4.7 μm, respectively. The design of bilayered structures was planned in such a way that the top layer can be used as entry for protein whereas low pore size bottom layer could not allow protein in this direction. The gold deposition on NAA was achieved by coating 10 nm thickness of gold film on top of the surface of NAA bilayer via sputtering technique. The greater sensitivity was attained by increasing the layer of gold film on sensor surface which increases the RI of NAA. Initially, sensor materials were incubated with proteins at quantity of 1 mg/mL in PBS buffer solution for 2 h at 5 °C to avoid the binding on surface of gold layer. Further, samples were thoroughly washed in PBS solution to remove any free contaminants. Thus, identical procedure was carried out for protein in PBS solution for biosensing applications. It was found that the protein adsorption occurred on the surface of NAA due to electrostatic forces between protein and sensor substrate. The allowed protein was detected at the top layer using change in optical thickness between two layers. The following equation has been used to detect the protein by change in optical thickness: that is,

\[ \Delta \text{EOT} = \text{EOT} \ (\text{after protein penetration}) - \text{EOT} \ (\text{before protein penetration}) \]  

(1)

where EOT is effective optical thickness.

The schematic diagram of fabrication and structure of Au-NAA bilayer and fast Fourier transform graph of the optical biosensing response of Au-NAA bilayers before and after the penetration of protein into sensor structures are shown in Figure 12. As shown in Figure 12(b), the peak intensity was very less in overall gold coated NAA bilayer with infiltration of protein due to adsorption of protein in layer 1. The increase in effective RI of layer 1 will decrease the RI of overall sensor substrate. The effect of optical thickness was increased in layers 1 and 3 due to the above reason, whereas less shift in layer 2 optical thickness was observed. The same results were obtained for Au-NAA bilayer without protein where higher intensified peaks were formed. The Au-NAA bilayer with the protein (BSA) has been sensed by intensified peaks of layers 1 and 3.

In similar work, Yeom et al. fabricated Au-NAA for real-time detection of antigen (C-reactive protein) [57]. The fabricated NAA have dimensions of 100 nm pore diameter and depths of pores were 4.9 μm. The Au-NAA materials were obtained by electron beam evaporation method. Here, gold layer coating on sensor material acts as high SPR compared to bare Ag or Au metals. The Au coated sensor substrate has produced LSPR and it plays vital role in accepting and immobilizing antibody on sensor surfaces. The immobilized antibody has been captured by Au-NAA material which was rinsed and kept for 24 h in solution of 11-mercapto-1-undecanol and EtOH. The unwanted antibody was removed by treating sensor surface with Tris-HCl solution instead of PBS solution to avoid the damage. The detection of antigen using Au-NAA works on LSPR principle in which the allowed antigens react with antibody to change RI of the sensor substrate. The biosensing mechanism has been predicted by change in the reflectivity of the sensor substrate by real-time biosensor detection method (Figure 13).

Similarly, Hiep et al. fabricated photonic structure of Au-NAA film for the detection of biomolecular interaction using interference localized surface plasmon resonance...
(iLSPR) system [58]. The gold coated NAA film (Au-NAA) has thickness of 270 nm which was rinsed in 11-amino-1-undecanethiol hydrochloride solution to functionalize with free amino groups available at substrate surface. A mixture of PBS and BSA solutions was used for rinsing NAA surface. The iLSPR system was used for the detection of binding interaction of two different biomolecules on NAA surface. The RI was slightly changed due to interaction of biotin-avidin binding on NAA surface and further increased for BSA binding with avidin on sensor surface. The limit of detection for avidin using Au-NAA film by iLSPR was found to be 100 ng/mL. Schematic diagram shows iLSPR biosensor and wavelength depends on reflectance spectra changes by biotin-avidin and biotin-BSA binding interaction on Au-NAA surface (Figure 14).

5.2. Detection of DNA. Kim et al. fabricated “cap like” gold structures which were coated on NAA for the detection of DNA hybridization sensing mechanism using LSPR microscopic system [59]. The fabricated NAA has the diameter and thickness of 60 nm and 5 μm, respectively. The immobilization of DNA on Au-NAA was achieved by mixing of 10 μM thiolated DNA probe and 100 μM 6-mercaptophexanol solutions which were further applied on sensor surface. A target DNA was hybridized with probe DNA on sensor surface by PBS solution. The sensing of probe DNA and target DNAs hybridization was captured by the increment in wavelength shifts (λ\text{max}) and relative reflected intensity which depends on change in the concentration of target DNA on the sensor surface. The saturation state was found at the concentration of target DNA which exceeded the limit of 10 μM. Schematic diagram shown in Figure 15 represents the working mechanism and sensing measurement of DNA hybridization on Au capped structure of NAA using LSPR system.

5.3. Detection of Cells. In biosensors field, researchers are trying to develop NAA with gold film coating for the detection of biomolecules such as proteins and DNA. However, less study has been carried out on the fabrication of Au-NAA for the detection of single living cell adhesion and seeding on sensor surfaces using SERS system. Toccafondi et al. fabricated biosensor substrate, that is, Au-NAA, for single cell sensing with the help of SERS [91]. Fabricated Al thin film on glass substrate has thickness of 500 nm by e-beam evaporator followed by anodization process. The prepared film structure had pore diameter and depth sizes of 50 and 120 nm, respectively. Thick gold layer of 25 nm on NAA was coated by e-beam evaporator and N2a cell lines were used from mouse neuroblastoma for living cell sensing application by bare NAA and Au-NAA using SERS system. In this experiment, 5 × 10^4 N2a cells were mixed in 2 mL of high glucose medium and cells were grown in this solution at 37°C with humidified CO_2 atmosphere. Later, cells were allowed for sensing application using bare NAA and Au-NAA. During the sensing process, different diameters (60 and 120 nm) of bare NAA were used to adhere to the N2a cells without using any adhesion solution which was due to biocompatibility and surface roughness of the material. The adhered cells were captured with fluorescent dyes by confocal fluorescence microscopy. However, the aim was to develop Au-NAA for living cell sensing applications using SERS. The excellent micro Raman scattering improvement has been achieved by cell seeded on Au-NAA. The sensing of Au-NAA material was done by laser light obtained from micro Raman spectroscopy. Enhancement in Raman peaks was observed in the presence of N2a cell due to adhesion and growth on surface of sensor material. On the other hand, Toccafondi et al. fabricated biocompatible Au-NAA for the detection of fibroblast living cells using SERS and fluorescence microscopy [92]. The SERS signals were high for the higher pore dimension. The cells were grown on Au-NAA surface due to biocompatibility of material. The living cells (NIH-3T3 cells) were incubated in Au-NAA surfaces for 2 days and grown on the surface. The better sensitivity was achieved for the bigger pores (130 nm).
and smaller pore lengths (50 nm) of Au-NAA for living cell sensing using SERS system.

5.4. Detection of Fluorescent Molecules. Das et al. developed novel Au-NAA with honeycomb pore structure for the biosensor applications with fluorescent molecules such as cresyl violet, rhodamine 6G, and green fluorescent protein-mut2 using SERS system [60]. Honeycomb pore structures of NAA with the pore size and wall thickness of 15 and 36 nm, respectively, were fabricated. Three different honeycomb pore structures were fabricated: NAA1 with pore size and wall thicknesses of 181 ± 45 and 82 ± 35 nm (bigger pores), NAA2 of 82 ± 7 and 21 ± 7 nm (medium pores), and NAA3 of 29 ± 4 and 13 ± 4 nm (smaller pores), respectively. The SERS signals were enhanced by coating 25 nm of gold film on NAA by thermal evaporation, namely, Au-NAA1 with pore size and wall thickness of 157 ± 35, 104 ± 35 nm, and Au-NAA2 with 59 ± 9, 40 ± 9 nm, and Au-NAA3 with 15 ± 3 and 36 ± 3 nm, respectively. To achieve the sensing, fluorescent molecules were deposited on different Au-NAA substrates by chemisorption method and tested with the molecules of concentration 10^6 M. Further, cresyl violet, an organic dye, was chosen which is routinely used in nerve tissue sector for biological stain. At first, cresyl violet molecules were coated on different Au-NAA structures revealing that SERS signals were enhanced due to NH₂ groups which were forcefully attached on sensor surfaces attributed to vibrational modes at 591 cm⁻¹. These molecules were strongly attached on surface of Au-NAA3 due to low porosity whereas in case of Au-NAA1 and Au-NAA2 molecules were not attached due to medium and high porosity, respectively. It was found that the enhancement in SERS measurement of Au-NAA2 compared to other two substrates due to low pore size and wall thickness. The enhancement was observed in SERS measurements for rhodamine 6G and green fluorescent protein-mut2 molecules which was due to the extensive
use of rhodamine 6G as a dye molecule in fluorescence microscopy. Due to the fact that molecular adsorption on metal surfaces is because of electronegativity of metals, this electronegativity is high for noble metal Au compared to Ag. Thus, the adsorption of molecules was enhanced by Au coating on sensor substrates. The rhodamine 6G molecules were deposited over surfaces of Au-NAA1 and Au-NAA2 to test in SERS systems. These molecules were strongly bound to Au surfaces and the enhancement in SERS was observed with strong intensity of vibrational modes at 1361 and 1510 cm$^{-1}$. The green fluorescence protein-mut2 was a fluorescent protein which has a capability of fluorescence recovery to use in slow biological processes. The SERS spectra enhanced through the fluorescent proteins which were coated on Au-NAA2 surfaces and SERS peaks were obtained at 1500 cm$^{-1}$. The schematic diagrams of honeycomb nanopore structure of Au-NAA and the SERS spectra are shown in Figure 16 which reveals the detection of (i) cresyl violet and (ii) rhodamine 6G using SERS.

5.5. Detection of Binding of Alkyl Thiol. Previously, several articles show developing of NAA for the detection of immobilized biomolecules on sensor surface by soaking the substrate in mixer of preferred buffer solutions like PBS/Tris-HCl containing biomolecules. However, biomolecules sometimes may not properly bind on the substrate with this technique. On the other hand, the biomolecules can also be immobilized onto the substrate by flowing the biomolecules taken in a mixture of buffer solutions within microfluidic channels for better binding due to higher chances of biomolecules coming in contact with the surfaces. The microfluidic system comprising sensor substrate has the advantages such as healthy binding biomolecule, less amount of sample for the detection, and rapid sensitivity in microchannels. For the first time Kumeria and Losic developed Au-NAA comprising microfluidic channel for label-free optical biosensing for the detection of binding of alkyl thiol on sensor substrate by enhancement of reflectivity in interferometric spectroscopy signals [61]. Pore size and pore depths of fabricated NAA structures were of 60 nm and 4 μm, respectively. In addition to that, scanning electron microscopy was used to measure the interpore distance and it was about 60–100 nm. The reflectance interferometric spectroscopy (RIS) signals were enhanced by the pores and surface area of internal pore structures of NAA. The fabricated controlled parameter of NAA has been coated by different metals like Au, Ag, Cr, Pt, and Ni by atomic layer deposition. However, Kumeria and Losic have not observed significant reflectance interferometric intensity signal originating from Pt, Ag, and Ni metals. The better interferometric signal has been obtained for Au and Cr metals due to enhancement of reflectivity on these thin metals. The optical biosensing performance was performed only by Au-NAA structure because Au metal has biocompatibility and chemical stability and is preferred for particular molecules binding detection for biosensing applications. The set-up comprises Au-NAA into microfluidic device with connected syringe pump and fiber optic probe. The optical biosensing showed by 1mM thiol molecules which were passed into Au-NAA substrate mounted microfluidic chip reveals higher wavelength with respect to change in effective optical thickness obtained for adsorption of molecules on Au surfaces. Furthermore, even at low concentration of molecules it has shown better sensitivity. The effective optical thickness has been increased with increase in the concentration of thiol molecules on sensor surfaces. They also analyzed real-time sensing performance of the molecule bounded with thiol group on sensor substrate which shows excellent maximum wavelength. The schematics shows the Au-NAA embedded
in microfluidic channel for detection of alkylthiol binding by optical biosensor using RIS system (Figure 17).

5.6. Detection of Circulating Tumor Cells. Kumeria et al. developed label-free microfluidic channel based biosensor on Au-NAA for the detection of circulating tumor cells (CTC) by reflectrometric interference spectroscopy [93–96]. Biotinylated anti-EpCAM was immobilized on the NAA surface through multiple chemical steps followed by integrating the porous NAA substrate into the microfluidic device. Spiked tumor cells in PBS and whole blood samples were injected into the device and captured with high efficiency by using interferometric spectroscopy. The fabricated NAA has pore diameter and lengths of $32 \pm 3$ nm and $4 \pm 0.2 \mu m$, respectively. The sensor surface was coated with ultra-thin Au layer of 8 nm by thermal vapor deposition technique. Multistep surface modification was adopted: Au-NAA were soaked in 10 mM solution of mercaptoundecanoic acid (MUA) in order to bind it covalently onto the sensor surface, followed by soaking in solution containing streptavidin molecules at 100 $\mu g/mL$ concentration where streptavidin binds to MUA. Further, the streptavidin functionalized substrate was incubated in 50 $\mu g/mL$ of anti-EpCAM antibody in PBS buffer solution. The PMMA based microfluidic chip was obtained by computer numerically controlled (CNC) micromaching method. This microfluidic chip was formed with two channels with sensing platform for Au coated NAA substrate on the glass substrate. The working principle of detection involves the use of white light emitted from optical fiber probe which was focused on anti-EpCAM modified sensor substrate incorporated in the chip and light was reflected back from the surface into the same optical fiber probe system. The change in effective optical thickness on sensor surface was attributed to the change in RI of the surface. The RIS signals were obtained after the flow of CTCs into the channels, due to binding of CTCs onto anti-EpCAM functionalized Au-NAA substrate. The effective optical thickness was increased due to continuous binding of CTCs onto anti-EpCAM coated sensor surface. The interferometric signals were changed a bit due to unbound CTCs which were removed from the surface after washing. The detection limit of CTCs was found to be less than 1000 cells/mL, which is significant in practical
applications and clinical diagnosis. It was also observed that, when the flow rate of CTCs was higher, there was no notable increase in effective optical thickness whereas, when the flow rate was low, effective optical thickness was enhanced by slow ascend and continuing the number of CTCs increment on the sensor surface.

5.7. Optical Properties of Au-NAA. The optical properties of Au-NAA for the application in biosensing by using reflectance spectroscopy have been developed by Hernández-Eguía et al. [97]. NAA substrates were fabricated with various porosities in the range of 14 to 70%. The expected pore diameters were in the range 38.6 to 90 nm, on which 20 nm Au was coated by sputtering method. The prepared Au-NAA samples were analyzed by reflectance spectroscopy and Fabry-Perot interferences (FI). The interference fringes consist of NAA film and are surrounded by the incident medium of air and the substrates were observed. The FI depends on RI of the material; that is, FI was increased upon increase in RI of Au-NAA substrate, whereas, noncoated NAA samples showed decrease in FI due to less RI compared to Au-NAA. The study on controlled thickness of gold layer was performed to obtain better reflectance spectra in the infra-red region. Thus, by controlling the thickness of the gold layer on NAA surface the better performance of the material can be achieved in terms of sensitivity.

6. Aluminum Coated NAA (NAA/Al) Biosensor

6.1. Detection of Protein. Further, the improved label-free optical wave guide biosensor based NAA/Al multilayer film has been developed by Hotta et al. for the detection of BSA through optical wave guide spectroscopy [62]. The wave guide experiment was functioned by measuring the reflection spectra of multilayer thin films using Kretschmann configuration. The multilayer film was fitted with glass prism and xenon lamp was used to irradiate the light onto prism. The incident light was slightly adjusted on prism by polarizer. Then, the reflected light was collected in photonic multichannel analyzer. A polydimethylsiloxane (PDMS) flow cell has been used to flow the analyte solution on multilayer film sensor. The irradiated light was incident on analyte solution and reflected spectra have been used for detection of analyte. The different structures of NAA/Al multilayer films were obtained by varying the substrate thickness. Al films which have been coated on glass substrate with thickness of 180 nm by thermal evaporation and films were anodized with formation of 34% porosity, and obtained pore diameter and lengths were of 39 and 220 nm, respectively (film a). Similarly, the same deposition method was used to fabricate Al film of thickness 500 nm and it was anodized with formation of 45% porosity to get cylindrical pore structured film of diameter and lengths of at 33 and 670 nm, respectively (film b). For the adsorption of BSA on NAA/Al film structures, the films were rinsed in mixture of prostate specific antigen and BSA solutions. The adsorption of BSA has been analyzed with respect to change in reflectivity on NAA/Al multilayer films. The reflectivity increased due to higher RI by increasing the concentration of BSA on film b compared to film a surface. However, it was found that film b was highly red shifted for the adsorption of BSA due to higher surface area, high porosity, and larger pore length with pore size compared to film a. Thus, the sensitivity was more for multilayer structure of film b over film a. Schematic diagram and working principle of detection of BSA on sensor substrate using optical waveguide spectroscopy and graph showing binding on substrate versus concentration of BSA in the solution phase were shown in Figure 18. Similarly, Yamaguchi et al. fabricated NAA/Al multilayer thin film on glass substrate for detection of binding interaction of different molecules using optical wave guide sensor mechanism [98]. Thus, the sensitivity depends on change in the RI of sensor material where analyte molecules were immobilized on their surfaces. The optical wave guide
sensor got great attention as the small changes in RI of material can be detected; that is, the sensor could detect lower concentration of samples.

7. Platinum Coated NAA (Pt-NAA) Biosensor

7.1. Detection of Binding of Prostate Specific Antigen. An et al. developed Pt-NAA biosensor for the improvement of prostate specific antigen binding on sensor surface in interferometric method through pore widening [99]. Earlier same group reported pore widening method on porous Si where less sensitivity was observed [100]. However, the experiment of Pt-NAA biosensor has improved the sensitivity through fabrication of pore widening structure on NAA. The Pt-NAA was silanized by APTES solution and later this substrate was treated with Prolinker-A solution for binding of biomolecules onto the surface. The Prolinker-A modified sensor surface was bound with prostate specific antigen. The detection of antigen by pore widening on sensor surface by the prostate specific antigen solution of 100 ng/mL treated with Pt-NAA surface reveals that higher wavelength means rapid detection of binding of antigen molecules on sensor surface. The effective optical thickness has been increased with respect to pore widening on Pt-NAA biosensor surface which is due to larger antigen molecules occupied and bound to Prolinker-A on larger pore and the optical signals were enhanced to get rapid detection through interferometric spectroscopy. Overall, it was found that biosensing of NAA, Au-NAA, Al-NAA, and Pt-NAA showed better sensitivity towards targets analyzed using different spectroscopy techniques. In NAA, targets were immobilized on their surfaces via binding with antigen-antibody. However, NAA shows less sensitivity compared to other substrates which has been enhanced by shrinking the pore size of Al, Au, and Pt coated NAA. Further, Au-NAA was chosen for biosensing application due to their higher refractive index, surface plasmon, and biocompatibility compared to NAA, Al-NAA, and Pt-NAA. Au-NAA was able to detect biomolecules at very low concentration of 100 ng/mL. In the case of NAA and Al-NAA for the analysis of biomolecules, the concentration was set to be of 0.1/1 mg mL\(^{-1}\). The targets were immobilized by covalently binding with antigen-antibody on Au-NAA surface for rapid detection. Lastly, Pt-NAA have better sensitivity compared to Au-NAA but the cost effectiveness of platinum hinders the usage in commercialization purpose.

8. Conclusion

In this review, past and recent progress on surface modification and development of NAA platform by the anodization process for suitable biosensing applications have been discussed. By controlling the parameters such as applied voltage, selection of acids, anodization time, and temperature, the precise and highly porous structure of NAA substrates can be fabricated. The methods combined with analytical techniques provide great benefits in terms of technological aspects in biosensors field. Further, the sensitivity and better performance of NAA substrates can be enhanced by coating with gold nanoparticles and also by incorporating with microfluidic devices. However, some process to immobilize biomolecule on substrates requires selected buffer solutions, and some buffer solutions can damage the NAA substrate depending on the pH level, so the immobilization process may not occur on NAA surfaces which hinders the biosensing process. We believe that in future, the improvement of NAA sensor and modern NAA based sensing platforms will give better results. Further, this review reports sensing mechanism of various kinds of NAA to immobilized biomolecules at lower concentration which was examined by using several spectroscopic techniques.
Competing Interests
The authors declare that there is no conflict of interests regarding the publication of this paper.

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