Abstract
This work investigated the performance of overlapped gate-on-drain of a gate all around-tunnel field-effect transistor (GAA-TFET) biosensors by considering the dielectric modulated technique by immobilizing the targeted biomolecules in the cavity region curved under the overlapped gate-on-drain. The nanowire GAA-TFET device shows excellent controllability over the channel and reduces leakage current to a greater extent. Here, we tried to make the ambipolar nature of the TFET, an advantage for the biosensor by detecting the biomolecule using variation of ambipolar current of TFET. Due to structural arrangement, the nanocavity under the overlapped gate region suppresses the ambipolar drain current by increasing the dielectric constant of the targeted biomolecules. The device can show a variation of $10^2$ and $10^3$ amount of sensitivity for the variation of dielectric constant from 1 to 5 and, compared with the other TFET structure, the proposed overlapped gate-on-drain GAA-TFET biosensor shows higher sensitivity and low leakage with a highly controlled channel.

Keywords Ambipolar behavior · Biosensor · Dielectric Modulation (DM) · Tunnel Field-Effect Transistor (TFET) · Gate All Around-Tunnel Field-Effect Transistor (GAA-TFET) · Overlapping gate-on-drain

1 Introduction
At dawn, and evaluation of the first-ever developed biosensor gives everyone optimism that there is an alternative to traditional laboratory techniques for detecting biomolecules [1, 2]. The stupendous characteristics of biosensors created enormous interest to carry forward the research in this field of biosensors. The advancement in the technology helped to drift the biosensor field to the extreme level and spread their application to many areas like environment monitoring, food processing, and the public health care sector. The biosensor plays an essential role in the medical field because of the following characteristics: early detection, continuous monitoring, and disease diagnosis. The present corona (Covid-19) pandemic’s situation is the best indication of biosensor’s importance in medical diagnosis since the early identification of the disease can stop the virus’s outbreak transmission from one to many [1]. Therefore, with the biosensor’s aid, they can carry out tests, trace, isolate, and prevent form transmission of disease.

Based on the kind of techniques, such as bio-recognition and identification methodologies, the biosensors are classified. The biosensor has divided into two types based on the detection technique: (1) label-based detection and (2) label-free detection. Due to its simple and cost-effective design approaches, the label-free detection technique has gained tremendous popularity. The field-effect transistor (FET)-based biosensors created revolutionary change and development in the field of biosensors [2–7]. The electrical detection of the target biomolecules using FET by virtue of label-free detection can develop a biosensor with the following characteristics (1) high sensitivity, (2) high selectivity, (3) On-chip integration of both sensor and the amplifier (4) Cost-effective and (5) Possibility of mass production and Reusability. Many researchers were impressed and inspired by the FET-based biosensors’ characteristics, carried out immense research, and reported a good and fruitful development in FET-based biosensors. The performance of the FET-based biosensors is high in class, but still, they are beyond touching the tip of the sensitivity because of the following hectic issues (1) the theoretical constraint on the minimum achievable
 Transportation and tunneling of the charge carriers in TFET depend on two major factors; first, one is applied gate bias, and the second factor is how effectively the gate can control the channel. To inflate the gate controllability over the channel and enhance the tunneling rate, numerous techniques and gate engineering methods were introduced for TFET. The All-around gate is an engineered gate structure that yields excellent controllability over the intrinsic channel of TFET. The nanowire gate all around the TFET device induces superior controllability over the channel and reduce the leakage current [16, 17]. The size of the target biomolecules in the given analyte is very small, and the more the surface to volume ratio acquired by the sensor will exhibit more sensitivity for the detection of the target biomolecules. Therefore, the nanowire-based biosensor can result in high sensitivity compared to the bulk device due to the increased surface to volume ratio attained by the nanowire-based biosensors and this high density of the biomolecules over the nanomaterial result in the experience in the high gating effect. There is colossal research that reported that the nanomaterial-based (nanoparticles, nanowires and nanotubes) biosensor results in increased sensitivity compared to the bulk devices [28, 35–37].

Here in this work, we have investigated the performance of nanowire GAA-TFET-based biosensor by considering the overlapped gate-on-drain technique. The partial hybridization and the practical binding difficulties of biomolecules are essential factors for biosensors, and here, we have included these effects to evaluate the sensitivity of the GAA-TFET biosensor. The entire work was carried out in four sections, and in section II, we discussed the architecture and the device’s simulation strategies. Also, we elaborately described the device results and discussions in Section III by taking partial hybridization (PH) and practical binding difficulties (PBD) of the biomolecule into consideration and the conclusion followed in the subsequent section IV.

2 Architecture and simulation setup of GAA-TFET biosensor

Figure 1 represents the 3D schematic arrangement of the proposed nanocavity curved overlapped gate-on-drain GAA-TFET biosensor. The 2D cross-sectional view of the proposed GAA-TFET biosensor provides an insight view of the proposed biosensor’s spatial arrangement, which is illustrated in Fig. 2. The target biomolecules made immobilized in the nanocavity curved under the overlapped gate-on-drain oxide. A 1 nm thick oxide layer (SiO2) is used as an adhesion layer for the biomolecules immobilized in the nanocavity region under the overlapped gate-on-drain. This adhesion layer helps in strict binding of the biomolecules and immobilization in the nanocavity area. A single silicon
nanowire is considered as the substrate for the biosensor design, and the simulation parameters are listed in Table 1.

Figure 3 represents the simulation structure of the proposed GAA-TFET biosensor with immobilized biomolecules in the cavity region. The simulation of the device is carried out by introducing a material having a dielectric constant (\(K > 1\)) corresponding to the biomolecules (i.e., protein, biotin, and streptavidin). The design parameters used for the GAA-TFET biosensor in this work is as follows: the radius of the silicon nanowire is 10 nm, and the channel length is 70 nm (\(L_{ch}\)), drain length is 50 nm (\(L_{D}\)), and the source length is taken as the 50 nm (\(L_{S}\)). The source is doped with \(p\)-type (\(1 \times 10^{19} \text{ cm}^{-3}\)), the drain is with \(n\)-type (\(5 \times 10^{18} \text{ cm}^{-3}\)), and the channel is doped with \(6 \times 10^{15} \text{ cm}^{-3}\). The work function (\(\Phi\)) for the gate electrode is selected as 4.8, and the thickness of the cavity is taken as 4 nm (\(T_{cavity}\)), and the overlapped-drain on gate length is taken as 50 nm (\(L_{ov}\)). Table 1 represents the device parameters used for the simulation.

The device’s simulation is performed by assuming that the cavity is filled by biomolecules entirely and made immobilized in the embedded nanocavity. The dielectric constant of the biomolecule in the overlapped gate-on-drain cavity is varied from \(K = 1\) to \(K = 10\) to investigate device performance under all circumstances. The benchmark reference value for measuring the sensitivity is competed filling the cavity with air having a dielectric constant value equal to (\(K = 1\)). The practical binding difficulties (PBD) and the partial hybridization (PH) effects of the biomolecules are taken into account to investigate the device’s performance [19–22].

SILVACO Atlas, a commercially available TCAD simulator, is used for simulating the device structures [18]. The non-local BTBT (band-to-band tunneling) model is preferred for computing the tunneling probability of charge carrier at the interface; the Auger model, Shockley–Read–Hall recombination/generation model, Kane’s model, the bandgap narrowing (BGN), the Fermi–Dirac carrier distribution models with default parameters are taken into consideration while doing the simulation of the device. The fabrication of the proposed device is complex compared to the conventional TFET devices, and the improved technology made the 3D device fabrication simple and feasible. Numerous researchers give many techniques to overcome these difficulties in device fabrication, and they are listed in works of literature with the modern nanolithography technology, etching and epitaxial growth techniques, which are fabricated into a real-time device [33–35].

### 3 Results and discussions

In this study, the ambipolar characteristics of the TFET are investigated immensely because the change in the ambipolar current (drain current \(I_{d(amb)}\)) of the TFET is taken
as the sensing (s) parameter for detecting the presence of the target biomolecule. At first, we have made few assumptions before investigating the GAA-TFET biosensor that the nanogap under the overlapped gate on the drain is filled with biomolecules (i.e., protein, biotin, streptavidin and Uri case) and immobilized in the cavity. The dielectric constant of the immobilized biomolecules inside the cavity is varied in a range from $K = 1$ to $K = 10$, and the corresponding response in the ambipolar characteristics of the device noted down to analyze the device sensitivity as the dielectric constant of the biomolecule changes inside the nanocavity region results in the energy band bending near the drain-channel junction. Figure 4a shows the variation of the energy band diagram of the GAA-TFET biosensor in counter to the change in the dielectric constant of the biomolecule. Here, we changed the dielectric constant of the biomolecule form $K = 1,2,3,5,7,10$, and it is observed from the corresponding energy band diagram that as the dielectric constant of the biomolecule is increases, it results in an increment in the bandgap at the channel-drain junction. The increment in the bandgap is linear up to $K = 5$, and after the bandgap remains constant and, this happened because of the limitation of the potential inside the nanowire GAA-TFET biosensor. If the dielectric constant value of the biomolecule is further increased beyond $K = 10$, the energy bandgap of the device remains stable due to fringing field effects given by surrounded gate structure.

From Fig. 4a, it is observed that the bandgap increases with increasing the dielectric constant of the biomolecule, and due to this increased energy bandgap, the tunneling width of the charge carrier at the channel-drain junction increases and results in the reduction of the ambipolar current of the device. The corresponding ambipolar current for the change in the dielectric constant of biomolecules is shown in Fig. 4b. The transfer characteristics of the overlapped gate-on-drain GAA-TFET biosensor for different dielectric constant values of the biomolecules ranging from $K = 1$ to $K = 5$ are shown in Fig. 4b. Here, we represented the absence of the biomolecule by taking a new material with dielectric constant $K = 1$ (Air), and this is taken as the reference for detecting the presence of biomolecule and measuring the device’s sensitivity. The mathematical formulation of the sensitivity of the device is expressed in Eq. 1.

\[
\text{Sensitivity } (S) = \frac{I_{ds(air)}}{I_{ds(bio)}}
\]  

(1)

For the dielectric modulation-based biosensor, the biomolecule’s polarity either improves or decreases the sensitivity for a low dielectric constant ($K$) biomolecules depends on the types of Si substrate doping where the cavity is created. Here, we considered the overlapped gate-on-drain technique where the sensitivity depends on the drain depletion and the higher the drain depletion, the higher the sensitivity. The charge of the biomolecule can show the notable impact on sensitivity for low dielectric constant valued biomolecules. However, when the dielectric constant of the biomolecule increases, it will dominate and reduce the charge of the biomolecule on sensitivity [23–27].

Figure 5a represents the proposed biosensor’s transfer characteristic for the dielectric constant values $K = 5, 7, 10$. From this Fig. 5a, we observed that even the dielectric properties of nanowire gate all around-TFET (GAA-TFET)-based biosensor with overlapped gate-on-drain with increasing dielectric constant from 1 to 5 and $V_{ds} = 0.5 \text{ V}$.
constant value is doubled, it shows a little variation in the ambipolar current, which is neglected and treated as constant. This is due to the effect of the energy band profile modulation (BPM), where the further widening of the bandgap is not possible because of the limitation on the minimum doping profile of the drain [27–31]. Even the surrounded gate structure shows high controllability over the channel, but the technique overlapping gate-on-drain depends on drain doping concentration. Here, we considered the drain doping of $5 \times 10^{18}$, which cannot further reduce due to the sensor’s construction constraints [27–31]. Figure 5b represents the potential variation of the GAA-TFET biosensor for different dielectric constant values of the biomolecules. The increase in the value of the dielectric constant of the biomolecule in the nanocavity influence the inter-trapped charge carriers near the channel-drain junction. These inter-trapped charge carriers create a potential change in the nanowire semiconductor, and as a result, the band starts changing their position.

The horizontal electric field along the device is shown in Fig. 6 for the targeted biomolecules with different dielectric constant values. Figure 6 shows that the overlapped gate-on-drain region possesses a high electric field near the channel-drain junction than the source-channel junction due to biomolecule presence under the cavity region of the overlapped gate-on-drain. This positive effect further increases the depletion width at the channel-drain junction and reduces ambipolar transport. The generalized expression for measuring the device’s sensitivity is expressed by Eq. 1, and the corresponding values are listed in Table 2. The measured sensitivity values from Table 2 shows that the proposed biosensor’s sensitivity increases linearly with the dielectric constant value up to $K = 5$ and, for higher dielectric values of biomolecules, i.e., $K > 5$, there is a slight downfall of sensitivity which is negligible and treated as constant. For the higher dielectric constant value of the biomolecules, the

\[
\text{Sensitivity (S)} = \frac{I_{ds(\text{air})}}{I_{ds(\text{bio})}}
\]
nanowire GAA-TFET shows no variation in the sensitivity because of the drain doping limitation [27–32]. The bandgap cannot increase further near the channel-drain junction even at the higher dielectric constant value of the biomolecules.

Figure 7 represents the drain current sensitivity plot of the nanowire GAA-TFET biosensor with overlapped gate-on-drain. The proposed biosensor can express the sensitivity, which is the ratio of drain current without biomolecule (i.e., \( k = 1 \)) to with the presence of biomolecule, with high sensitivity \( 10^6 \) for a wide range of biomolecules (i.e., \( K = 5 \)) and it maintained to all the higher dielectric constant values of the biomolecules.

The nanowire GAA-TFET biosensor shows excellent improvement in the device sensitivity in terms of high drain current variation. Apart from this, the nanowire GAA-TFET biosensor is capable of exhibiting high, prominent threshold voltage characteristics [22–32]. The device’s surrounded gate structure facilitates the gate with high controllability over the intrinsic channel and makes it operate at low voltages. Figure 8a represents the threshold voltage variation of the proposed nanowire GAA-TFET biosensor for different biomolecules. It is observed in Fig. 8a that the increment in the dielectric constant value of immobilized biomolecule inside the nanocavity reduces the threshold voltage of the device to a greater extent, and this is due to the increment of the overlapped gate capacitance near the channel-drain junction and reduced leakage current. We have taken the variation in the device’s threshold voltage as the parameter for detecting the biomolecule. The threshold voltage sensitivity is plotted in Fig. 8b using the following mathematical relation.

\[
\text{Sensitivity} \left( S_{V_{th}} \right) = \left( \frac{V_{th}(\text{air}) - V_{th}(\text{bio})}{V_{th}(\text{air})} \right)
\]

All results discussed in this section are performed by assuming that the nanogap region under the overlapped gate is filled completely with biomolecules. However, this case is impossible due to the biomolecules associated binding difficulties and the fill factor. Partial hybridization (PH) is one of the essential factors to be considered when designing a biosensor, and this effect occurs due to incomplete filling of the cavity and the steric hindrance effect of the biomolecules. Many people deliberately discussed these PH effects and the binding difficulties to realize the biosensor’s real-time performance [19–22].

Here we have incorporated the biomolecules’ binding difficulties by partial filling the cavity with air in the nanocavity.
region. Here we have taken three cases where the air can occupy the nanogap into different length, which is showed in Fig. 9. (b), i.e. 10 nm, 20 nm, and 30 nm. For all three cases, the partially filled biomolecules dielectric constant is taken as constant, i.e. \( K = 5 \), and we have noted down the respective drain current of the biosensor for all three cases, and the corresponding drain currents are plotted in the Fig. 10. Interestingly the PH and the binding problems associated with the biomolecules cannot show an impact on the device performance due to high controllability of the gate over the channel.

4 Conclusion

This paper presented a GAA-TFET-based biosensor using the dielectric modulation (DM) technique by placing the biomolecules inside the nanocavity curved in the oxide dielectric overlapped gate-on-drain region. Here, we used the nanowire GAA-TFET device for getting higher controllability over the channel, which yields higher sensitivity. It is observed that the device shows a variation of \( 10^3 \) amount of drain current \( (I_{ds(amb)}) \) by increasing the dielectric \( (K) \) constant value of the biomolecules from 1 to 5. It is also concluded that ambipolar transport, which is the drawback of the TFET for digital circuit applications, turns into an advantage for biosensor applications. The excellent and remarkable performance of the proposed GAA-TFET-based biosensor made it an advancement in the field of biosensors. In the future, this will play a vital role in the detection of biomolecules with controlled gate leakage current with superior low power functionality.

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Declarations

Conflict of interest The authors declare that they have no conflict of interest.

Consent for publication The authors have given Consent for Publication as per the journal policy.

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