CRISPR/Cas13a-Powered Electrochemical Microfluidic Biosensor for Nucleic Acid Amplification-Free miRNA Diagnostics

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Noncoding small RNAs, such as microRNAs, are becoming the biomarkers of choice for multiple diseases in clinical diagnostics. A dysregulation of these microRNAs can be associated with many different diseases, such as cancer, dementia, and cardiovascular conditions. The key for effective treatment is an accurate initial diagnosis at an early stage, improving the patient’s survival chances. In this work, the first clustered regularly interspaced short palindromic repeats (CRISPR)/Cas13a-powered microfluidic, integrated electrochemical biosensor for the on-site detection of microRNAs is introduced. Through this unique combination, the quantification of the potential tumor markers microRNA miR-19b and miR-20a is realized without any nucleic acid amplification. With a readout time of 9 min and an overall process time of less than 4 h, a limit of detection of 10 pm is achieved, using a measuring volume of less than 0.6 µL. Furthermore, the feasibility of the biosensor platform to detect miR-19b in serum samples of children, suffering from brain cancer, is demonstrated. The validation of the obtained results with a standard quantitative real-time polymerase chain reaction method shows the ability of the electrochemical CRISPR-powered system to be a low-cost, easily scalable, and target amplification-free tool for nucleic acid based diagnostics.

MicroRNAs (miRNAs) are composed of 18–25 nucleotides in their mature form and play an essential role as regulators of gene expression in many biological processes by binding to specific messenger RNA targets and promoting their degradation or translational inhibition.[1,2] In recent years, the popularity of miRNAs in research has been constantly growing, as the presence or dysregulation of distinct miRNAs can indicate specific medical conditions. For instance, in cancer research the up or downregulation of miRNA expression levels has been linked to certain cancer types, including lung cancer or brain tumors. This offers new possibilities for the detection and monitoring of such diseases, which is why miRNAs will continue to come into the focus of clinical diagnostics.[3–5]

Traditional methods of studying miRNA levels include microarrays, RNA sequencing methods and, as the current gold standard, the quantitative polymerase chain reaction (qPCR).[6] Each of these methods, despite being powerful, have their limitations. Bulky and expensive equipment, intensive sample preparation, or long turnaround times limit most of them to well-equipped laboratories, restricting such screenings to developed countries. Therefore, there is a pressing need to develop easy-to-use, portable, and amplification-free methods for miRNA detection at the point of care (POC). This will allow a fast, versatile, and low-cost
quantification of miRNAs not only in the western world but also in resource-limited regions, where people do not have access to well-equipped laboratories and where cost efficiency is even more important.

Today, there are a few reports tackling these challenges, for example, by improving current detection methods, using electrochemical assays\(^\text{[7,8]}\) or developing pH-based POC tests.\(^\text{[9]}\) However, these approaches still need error-prone amplification steps prior to the miRNA detection, have rather tedious preparation steps, or are limited by the design of suitable primers.\(^\text{[10,11]}\) To address these issues, we combine microfluidics with an electrochemical signal readout to develop a sensitive (i.e., target amplification-free) and selective diagnostic test, while enabling a miniaturization for POC testing.\(^\text{[12,13]}\) We further apply the newly discovered CRISPR/Cas13a technology, which is able of targeting almost any RNA, to our developed electrochemical biosensor, creating a novel and powerful tool for miRNA diagnostics (Figure 1a).

In the recent past, studies, employing CRISPR-associated methods for the detection of nucleic acids, have been utilized using different CRISPR-associated (Cas) effectors.\(^\text{[14]}\) Varying Cas effectors were used for targeting different nucleic acids, like the Cas9 effector for the detection of double-stranded DNA (dsDNA),\(^\text{[15–17]}\) along with the Cas12 for the detection of single-stranded DNA (ssDNA).\(^\text{[18–21]}\) The microbial CRISPR effector Cas13a (previously named C2c2), displays, in contrast to the other Cas effectors, a triggered cleavage capability of nontarget single-stranded RNAs (ssRNAs) in the surrounding.\(^\text{[22,23]}\) Taking advantage of this characteristic, we replace synthetic nucleic acid amplification steps by a Cas13a-driven signal amplification. The specificity of the detection is given by the need of a target-specific CRISPR RNA (crRNA) that guides the Cas13a to the RNA sequence of interest (Figure 1b).\(^\text{[23,24]}\) Upon recognition of the complementary RNA sequence, the cleavage ability of the enzyme is activated. The addition and subsequent cleavage of a reporter RNA (reRNA) can thus be used for a quantitative readout of RNA levels.\(^\text{[25]}\) Since after the activation of the Cas/crRNA complex the collateral cleavage mechanism is, in contrast to other Cas effectors, an ongoing process, the amount of cleaved reRNA is a time-depending progress, resulting in self-amplification. These two specific characteristics of the Cas enzyme enable a highly selective and sensitive detection. So far, Cas13a was used for the optical detection of virus RNAs or even for the detection of plant genes,\(^\text{[26]}\) whereas no research work exists up to now, combining CRISPR/Cas13a with an electrochemical detection method for miRNA diagnostics.\(^\text{[22–25,27–30]}\) Here, we present the CRISPR technology on a microfluidic electrochemical biosensor for measuring miRNA levels of the potential brain tumor marker miR-19b in serum samples from patients, suffering from brain cancer.

For the detection of miRNAs, a low-cost and easy-to-use electrochemical biosensor was manufactured, using the DFR technology.\(^\text{[31,32]}\) By stacking multiple developed DFR foils onto a platinum patterned polyimide substrate, the microchannel and the electrodes were realized. The microfluidic channel consists thereby out of two distinct sections: (i) an immobilization area and (ii) an electrochemical cell, separated by a hydrophobic stopping barrier (Figure 1c). The immobilization area was filled and functionalized through an inlet by capillary forces, until the flow reaches the stopping barrier. This allows an easy handling of the chip, combined with an automatic metering (<0.6 µL) of the introduced biomolecules and, in addition, prohibits a contamination of the electrodes. In the electrochemical cell,
the detection of enzymatically produced hydrogen peroxide ($\text{H}_2\text{O}_2$) takes place, using a three-electrode setup, comprising a platinum working and counter electrode together with a silver/silver chloride reference electrode.

For the CRISPR/Cas13a-powered miRNA detection, the surface of the immobilization area of the biosensor was prefuntionalized by applying streptavidin to the chip inlet, which is followed by a blocking step, using bovine serum albumin (BSA) (1% in 10 mM PBS). After each incubation step, a washing step is performed, where the microchannel is flushed with wash buffer (50 µL of 0.05% TWEEN 20 in 10 mM PBS). By applying a vacuum to the channel inlet, all unbound biomolecules are removed, without contaminating the electrochemical measurement cell. For the activation of the Cas13a and the subsequent cleavage process, the enzyme Cas13a was mixed in a standard microcentrifuge tube with its target-specific crRNA, a biotin and 6-FAM (6-fluorescein amidite) labeled reRNA, and the sample of interest, potentially containing the target miRNA. The mixture was incubated in the standard tube for 1 to 24 h at 37 °C, where the Cas13a forms a complex with the target-specific crRNA. In the presence of target miRNAs, the Cas13a gets activated, resulting in a collateral cleavage of the surrounding reRNA.

After the off-chip targeting of the miRNAs, the mixture was subsequently applied to the prefuntionalized microfluidic chip, where the cleaved and noncleaved reRNAs, depending on the quantity of active enzymes, bind to the immobilized streptavidin. Following a washing step, anti-fluorescein antibodies coupled to glucose oxidase (GOx) (0.83 µg µL$^{-1}$ in 10 mM PBS), which are only capable of binding to the uncleaved reRNAs, were introduced, enabling an enzymatic reading of the assay (Figure 1a).

For the assay readout, the biosensor chip was placed into a custom-made holder, allowing the electrical connection to the potentiostat and the fluidic connection to a syringe pump. By pumping a glucose solution (40 mM glucose in 10 mM PBS) through the microfluidic biosensor, GOx catalyzes its substrate, producing $\text{H}_2\text{O}_2$, which is amperometrically detected in the electrochemical cell. For a further signal amplification, a fully automated so-called stop-flow protocol was used.\[33\] Herein, the pump was programmed so that the flow of the glucose solution is stopped for a certain period of time, where $\text{H}_2\text{O}_2$ is produced by the enzyme and accumulates in the immobilization area. The pump automatically restarts the flow, the enriched $\text{H}_2\text{O}_2$ concentration is flushed over the working electrode, which results in a current peak with a specific peak charge. The obtained amperometric signal is directly proportional to the amount of immobilized GOx, bound to the uncleaved reRNA and, therefore, inversely proportional to the concentration of target miRNA in the sample.

The assay’s sensitivity was improved through optimizing each assay component in terms of incubation time and concentration (Figures S2–S10, Supporting Information). Furthermore, to overcome the diffusion limitation of the reagents, a dynamic cleavage process, through shaking the mixture solution, while incubating at 37 °C, was implemented. As it became apparent, the shaking of the solution hindered the catalytic activity of the Cas13a (Figure S11, Supporting Information), why on a static cleavage process was chosen. As the crRNA needs to bind to the Cas effector in order to initiate the cleavage activity, the shaking might hamper this binding process or even promotes the loss of already bound crRNA.

By preparing solutions, containing different concentrations (100 fm to 10 nm) of the miRNA miR-19b and miR-20a, incubating them at 37 °C and applying them to the functionalized channel, different calibration curves were recorded (Figure 2, Figures S15 and S16, Supporting Information). By fitting the measured data points to a four or five-parametric sigmoidal curve, a LOD of 10 pm equivalent to an amount of 500 amol of miRNA, was achieved for the miR-19b within 3 h incubation.

![Figure 2](image-url)
time and an overall interassay coefficient of variation of less than 10% (Figure 2b). As the cleavage of the reporter RNA is a time-dependent process, the prolongation of the incubation increases the amount of cleaved RNA and, therefore, reduces the amperometric signal (Figure S14, Supporting Information). To decrease the limit of detection furthermore, a calibration curve with an extended incubation time of the mixture solution for 7 h (Figure S15, Supporting Information) and 24 h is recorded (Figure 2c). Contraintuitive, we could not observe an improvement of the LOD for a 7 h cleavage time. In our opinion, this is due to the very low concentration of the target miRNA and the therewith resulting lack of active Cas13a enzymes in the solution. This leads to an unsatisfactory total catalytic activity of the effector with a very low reaction rate, which is not sufficient for a further significant signal reduction at very low concentrations of the target miRNA within 7 h of incubation. Comparing the different cleavage times in terms of sensor performance, a minimum LOD of 2 pM was achieved after a 24 h incubation, while a maximum dynamic range of roughly two orders of magnitude can be seen, at a cleavage time of only 1 h, along with a LOD of 18 pM (Figure 2a). In general, the longer the cleavage time, the lower the limit of detection and the lower the dynamic range of the resulting calibration curve (Table S6, Supporting Information). With that, our CRISPR/Cas13a-powered biosensor is able to reach the stated LOD of 10 pm after 3 h, showing its capability to detect picomolar concentrations of miRNAs, without any preamplification procedure of the target miRNA, while consuming reagent volumes less than 0.6 µL per incubation.

In order to validate the feasibility of our sensor concept for the detection of low miRNA concentrations in clinically relevant specimens, four serum samples of children, suffering from medulloblastoma, were tested. Medulloblastoma is an aggressive embryonal tumor of the cerebellum/forth ventricle, which is characterized by an inhibition of apoptosis and an increased proliferation, characteristics caused by altered expression patterns in miRNAs. It has been reported that, compared to healthy children, the concentration of the miR-17–92 cluster family, containing the here employed biomarkers miR-19b and miR-20a, are upregulated for this tumor type. Therefore, samples of patients, showing different clinical conditions, were chosen. Patients 1 and 3 achieved a complete remission of the tumor, whereas patients 2 and 4 developed a progressive state within a few months, which suggests that the patients 2 and 4 should have a significantly elevated miR-19b level. For the biochip measurements, the total RNA from the serum samples was isolated, using a commonly available RNA purification kit. The purified RNA was mixed with the Cas13a, its crRNA, specific to the miR-19b, and incubated at 37 °C. Consequently, the on-chip incubation and signal readout were performed as illustrated in Figure 1c.

To confirm our biochip measurements, the standard qRT-PCR method was executed to gauge the miRNA levels of the miR-19b as well. The measurements were performed from the same samples with a similar RNA purification technique. A serum medley of a pool of 20 healthy patients was used as a control measurement and treated the same way as the samples of the four patients, suffering from medulloblastoma. The results of both measurement methods show a good agreement within all four patient samples, especially by taking the nonlinearity of our measurement method and the fact that the qRT-PCR cannot distinguish between miR-19a and 19b into consideration (Figure 3a). Besides being a powerful method for the detection of nucleic acids, qRT-PCR-based methods normally require sophisticated systems and sample preparation, like for the reverse transcription of the complementary DNA (cDNA), a long hands-on time and well-trained operators. In comparison with that, our approach for miRNA detection simplifies the

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**Figure 3.** CRISPR-based off-chip miRNA targeting. a) Validation of our measurement principle, using four serum samples from children, suffering from medulloblastoma and a control sample of a pool of 20 healthy patients. For the biosensor measurements, error bars represent ± SD of n = 4 replicates; error bars of the qPCR measurement represent ± SEM of n = 3 measurements; two-tailed Student's t-test; ***P < 0.001; ****P < 0.0001. b) Schematic of the crRNA target region and the target sequences used for the detection. The mismatches in the target sequences are highlighted in red. c) Comparison of the cleavage efficiency of different miRNA targets, completely matching the complementary crRNA sequence (miR-19b), having a single mismatch (miR-19a) or do not match to a greater extend (miR-197). Applied concentration of 1 nM for each miRNA with n = 8 replicates, two-tailed Student’s t-test; ****P < 0.0001; error bars represent ± SD.
procedure and enables thereby the detection in a low-cost and easy manner, without the use of complex signal amplification methods or specially trained personnel.

For an accurate diagnosis, a highly selective detection of the target miRNA is crucial. To investigate the biosensor’s specificity, target miRNAs, differing from the miR-19b by a single nucleotide (miR-19a) or several nucleotides (miR-197), were tested (Figure 3b). For this, concentrations of 1 nM of the miR-19b, miR-19a, miR-197 as well as a mixture of the miR-197 and miR-19b were added to the sample solution, containing the Cas13a and the crRNA, complementary to the miR-19b. For the miR-197, showing poly-base mismatches, the Cas13a does not show any cleavage ability, as the hybridization of the target miRNA to the crRNA is not sufficient to activate the enzyme’s catalytic activity. Comparing the miRNA miR-19b with the miR-19a, containing a single-base variation in the middle of the sequence, the biosensor enables a single-base mismatch detection within a significance greater of P < 0.00001 at low concentrations (Figure 3c). As both miRNAs (miR-19a and 19b), together with the miRNAs miR-17, miR-18a, miR-20a, and miR-92a, are part of the miR-17–92 cluster family,[16] upregulated in patients, suffering from medulloblastoma, the exact distinction of these miRNAs is not compulsory necessary for an accurate diagnosis of this type of tumor. Additionally, to enhance the specificity of a future system, synthetic mismatches within the crRNA can further increase the signal, while the Cas13a is able to distinguish between single-base mismatches, as reported by Gootenberg and colleagues.[27] This is not only the case for single-base mismatches in the middle of the sequence but also for different mismatch locations or double mismatches, as extensively studied in literature for the CRISPR effector Cas13a.[23,29]

For a more versatile and easy-to-handle biosensor, we also envision a “one-for-all” sensor chip, containing a complete pre-immobilized assay. In contrast to the off-chip miRNA targeting, where the activation of the Cas13a and the cleavage process were performed off-chip in a standard tube, the “one-for-all” sensor chip allows the activation and the cleavage to be done on-chip, while incubating. For the assay, an anti-biotin antibody instead of streptavidin is used as a surface coating for the binding of the biotin-labeled reRNA. By using streptavidin to couple the reRNA to the surface for the on-chip cleavage process, we believe that the Cas13a was sterically hindered, resulting in an inefficient cleavage process of the immobilized reRNA (Figure S18, Supporting Information). To overcome this issue, a polyclonal anti-biotin antibody is used as a replacement for streptavidin, creating a greater distance to the channel surface and thus providing a better accessibility of the Cas13a for the cleavage of the reRNA (Figure 4a).

All assay components, including anti-biotin antibodies for the surface coating, the biotin and 6-FAM-labeled reRNA, and the GOx-labeled anti-fluorescein antibody, are immobilized in the biosensor’s immobilization area. For the detection of any miRNA, only the mixture of the sample, potentially containing the target miRNA and the target-specific crRNA/Cas13a complex need to be introduced to the microchannel and incubated at 37 °C. While the mixture is incubated, the enzyme cleaves, in the presence of any target miRNA, the immobilized reRNA and thus remobilizes the GOx-conjugated antibodies. Following a washing step to remove all unbound GOx-labeled antibodies, the assay readout can be performed by introducing the glucose solution (Figure 4a). The produced H2O2 results, as before, in a current density proportional to the amount of bound GOx and is reverse proportional to the concentration of target miRNA in the sample solution. For further information and visualization of the “one-for-all” sensor chip handling and cleavage process, we refer to Movie S1 (Supporting Information).

For a first examination of the on-chip detection with the “one-for-all” biosensor, a calibration curve was performed. Solutions, containing different concentrations (1 pM to 1 nM) of the miRNA miR-19b, were prepared and introduced to the completely functionalyzed microchannel and incubated at 37 °C for 3 h. Furthermore, to increase the cleavage capacity of the Cas13a, the solution is exchanged hourly during the incubation process. After the amperometric readout, the data points were fitted to a four-parametric logistic curve, resulting in a LOD of 2.2 nM with an overall inter assay coefficient of variation of less than 15% (Figure 4b). This proves the feasibility to transfer the complete assay to the biosensor for an on-chip detection of low amounts (<5 fmol) of any miRNA at the point of need in a low-cost manner, without the need of expensive equipment. Nevertheless, as the gap in the LOD values of the off-chip miRNA targeting versus the on-chip detection is evident, the on-chip detection needs further investigation and optimization.

Since the discovery of the CRISPR class 2 effector Cas13a in 2015,[14] different approaches for the utilization of the effector for biosensing applications were presented in literature.[27–29,30] However, no work has been reported so far for a CRISPR-powered microfluidic assay, along with an electrochemical detection method. Here, we have introduced the unique fusion of a microfluidic, electrochemical biosensor with the powerful CRISPR/Cas13a technology for the quantification of low concentrations of miRNAs in serum specimens. Electrochemical, microfluidic biosensors have shown in the past their capability of detecting low concentrations of analytes in clinically relevant samples on an easy-to-fabricate integrated sensing platform, without the need of extensive instrumentations.[39] In particular, we use the CRISPR/Cas13a technology for self-amplification upon recognition of the target miRNA, combined with a fully automated microfluidic stop-flow protocol for a further amplification of the electrochemical signal. By this extraordinary combination, our detection method requires no nucleic acid amplification prior to the detection procedure and, therefore, no additional costly reagents or specialist equipment, including trained personnel is needed.

Moreover, until now amplification-free CRISPR-powered biosensing methods are rare and only one approach, employing Cas9, exists for an amplification-free detection of dsDNA, based on a graphene field-effect transistor (gFET).[16] For the detection of dsDNAs, several incubation steps lead to the final functionalized CRISPR-Chip, having a readout time of down to 15 min, compared to our stop-flow amplified readout process of roughly 9 min (Figure S17, Supporting Information). The CRISPR-Chip aims for the detection of gene mutations for the diagnostic of Duchenne muscular dystrophy associated mutations in the human genome.[40,41] For this, they are able to use the whole human genome (1.9 pg mol−1), which leads to an amplification-free detection limit of 1.7 fmol (3.3 ng mL−1). By
The on-chip miRNA detection: a “one-for-all” biosensor. a) Schematic illustrating the on-chip cleavage procedure for samples with or without target miRNAs (blue). On the left, the sensor is depicted, showing the immobilization area, highlighted in yellow with the pre-immobilized assay, comprising an anti-biotin antibody surface coating along with a BSA blocking. The biotin and 6-FAM-labeled reRNA is immobilized to the anti-biotin antibody, where the GOx-labeled anti-fluorescein antibody binds to the reRNA. On the right, a mixture, containing the crRNA/Cas13a complex and a sample with and without target miRNAs (blue), is introduced into the biosensor to the completely pre-immobilized assay. The target activated enzyme cleaves the bound reRNA and thus enables the removal of the GOx-labeled antibody, resulting in a reduced amperometric signal. b) Calibration curve of the “one-for-all” biosensor for an on-chip miRNA detection, using the miR-19b as a target miRNA. The results are fitted with a four-parametric logistic fit, gaining a limit of detection of 2.2 nM with an interassay coefficient of variation of below 15%. With $n = 4$ replicates, error bars represent ± SD.

Figure 4. The on-chip miRNA detection: a “one-for-all” biosensor. a) Schematic illustrating the on-chip cleavage procedure for samples with or without target miRNAs (blue). On the left, the sensor is depicted, showing the immobilization area, highlighted in yellow with the pre-immobilized assay, comprising an anti-biotin antibody surface coating along with a BSA blocking. The biotin and 6-FAM-labeled reRNA is immobilized to the anti-biotin antibody, where the GOx-labeled anti-fluorescein antibody binds to the reRNA. On the right, a mixture, containing the crRNA/Cas13a complex and a sample with and without target miRNAs (blue), is introduced into the biosensor to the completely pre-immobilized assay. The target activated enzyme cleaves the bound reRNA and thus enables the removal of the GOx-labeled antibody, resulting in a reduced amperometric signal. b) Calibration curve of the “one-for-all” biosensor for an on-chip miRNA detection, using the miR-19b as a target miRNA. The results are fitted with a four-parametric logistic fit, gaining a limit of detection of 2.2 nM with an interassay coefficient of variation of below 15%. With $n = 4$ replicates, error bars represent ± SD.

Taking the molecular weight of our employed miRNA miR-19b (7.3 × 10$^3$ g mol$^{-1}$) for a similar calculation, a limit of detection (LOD) of our biosensor of 73 fg µL$^{-1}$ is achieved. For extending the CRISPR-Chip to detect ssRNAs, the effector Cas13a could be used as well. As the signal generation of the gFET is based on the amount of adsorbed and interacting charged molecules, the LOD of the CRISPR-Chip of 3.3 ng µL$^{-1}$ would most likely stay the same, leading to a LOD of roughly 0.45 µM for the here employed miRNA miR-19b.

One major drawback of many reported diagnostic tools for miRNA detection is the extensive preparation of patient samples for the quantification of the target analyte. Often isolation and purification kits are needed, to extract the desired target without any other interfering biomolecules. Even though, our electrochemical biosensor needs one kit as well for the RNA isolation from serum, it is a commonly available one with a wide range of application fields. In general, the abandonment of such purification kits would drastically improve CRISPR-based detection methods, allowing an easier handling and a faster gauging of the desired analyte.

Despite the powerful detection of other CRISPR-based detection methods, the employed chips are often highly complex and require multiple fabrication steps. The employed gFET for the CRISPR-Chip, for example, is produced by using silicon fabrication techniques, which normally requires sensible clean room stages, making it a powerful, but costly detection tool. Our employed biosensor is based on polyimide and dry film photoresist (DFR) layers, which are fairly easy to fabricate. On the other side, one clean room step is also needed for realizing the platinum electrodes of the electrochemical cell of our biosensor, which accounts for roughly two-thirds of the total fabrication costs. The employed reagents, like antibodies, are partly costly as well, resulting in an allover price of roughly 0.75 € per functionalized biosensor, manufactured under research conditions (Table S4, Supporting Information). In general, our sensor device or silicon-based systems can be produced in standard industrial facilities, which will reduce the manufacturing price per chip drastically.

In this work, we have shown that our CRISPR/Cas13a-powered biosensor efficiently and accurately detects miRNAs in small sample volumes, without the need of any nucleic acid amplification step. The clinical potential of our system was successfully evaluated by measuring different serum samples of pediatric patients, suffering from medulloblastoma in different stages. By examining the same samples with a standard quantitative real-time polymerase chain reaction (qRT-PCR) method, the findings of our biosensing chip were validated within a good agreement. These results indicate that our biosensor chip is capable of diagnosing miRNA-related diseases, like brain tumors, by bypassing of any preamplification step. We furthermore overcome limitations, including the restrictions of common primer designs, extending this technique to short target miRNAs previously difficult to detect and with the obtained minimal detection limit of 2 ps, we are able to measure miRNAs in the clinical range of circulating miRNAs in serum.[42]

Although we are rivaling other Cas13a-powered optical detection methods for longer RNAs (i.e., SHERLOCK) by a factor of 5[27] prior to their nucleic acid amplification step, as reported by Li and colleagues,[24] other electrochemical methods for the detection of miRNAs exist, having lower LODs as our presented approach.[43–51] The basic working principle of most of these electrochemical sensors is the immobilization of a (labeled) nucleic acid sequence, complementary to the target sequence, onto a screen-printed (gold) electrode, which enables an amplification of the signal, resulting in low LODs. Besides the higher material values used in these sensors, this also indicates a much higher hands-on time for the incubation process for the functionalization of the electrode, as each electrochemical chip must be target-specifically pretreated before the desired miRNA can be detected. In contrast to that, our biosensor chip can be easily prefuctionalized and stored for up to nine months.
under refrigerated conditions (Figure S13, Supporting Information) without the need of any target specific pretreatment prior to the detection of the desired miRNA, compared to other electrochemical systems, having a stability in the range of days or weeks.[54]

By presenting the possibility of transferring the complete assay for the miRNA detection on our “one-for-all” biosensor chip, we have shown the feasibility to detect any desired miRNA, requiring only the input of the Cas13a/crRNA–target mixture. Despite, the “one-for-all” biosensor needs further improvements in terms of sensitivity and reproducibility, it does show the practicability of this system for an easy-to-use setup, reducing many incubation steps, while decreasing the sample-to-result time. Overall, the unique fusion of the CRISPR/Cas13a technology with our microfluidic biosensor, combined with an electrochemical readout, allows the detection of the desired miRNA with high sensitivity and selectivity with a short hands-on time. This pioneers the way for a new generation of on-site miRNA diagnostic tools.

Future work will be the further reduction of the LOD by implementing different strategies, like an immobilization of the Cas/crRNA complex. With that, the target sample can be flushed over the immobilized complex, while the target sequence will bind to the complex and activates the catalytic cleavage mechanism of the Cas effector. Employing this strategy to the developed biosensor, a strong reduction in the LOD should be noticeable. Besides that, an additional electrochemical signal amplification via a so-called redox cycling by using interdigitated electrode arrays (IDAs) can be combined with our microfluidic stop-flow measurement protocol. In our previous published works, signal amplifications of more than 150 times are achieved, employing nanogap IDAs with 100 nm gap size.[55–57] Combining this technique with our CRISPR/Cas13a-based biosensor, the LOD of our system could be reduced to less than 100 fm.

Furthermore, a future multiplexed version of our biosensor, comprising several incubation channels, could allow the detection of multiple miRNAs from one single sample, resulting in a more accurate diagnosis of a certain disease.[58]

This will not only allow us to detect one specific miRNA but also enables the realization of a positive (by the use of a specific target concentration) and negative (employing no target) built-in control for each measurement. Such a CRISPR/Cas-powered multiplexed biosensor, being able of measuring up to eight miRNAs (Figure S20, Supporting Information), will also deliver a more complete picture of the patient’s disease, for which nowadays expensive and lab-intensive equipment is needed.

Experimental Section

All experiments and methods are described in detail in the Supporting Information.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.
