## Minor groove binder probe real-time RT-PCR for detection of footand mouth disease virus in Egypt

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#### Abstract

Shorter, more specific minor groove binders (MGBs) probes are dsDNA-binding agents attached to the 3' end of TaqMan probes that could be designed strictly to invariant region. Application and assessing of a new trend for viral detection in Egypt depending on MGB probe real-time RT-PCR (rRT-PCR) was applied on local FMDV serotypes O, A, and SAT2. Moreover, FMDV O were detected using two serotype specific primer sets by SYBR Green real-time RT-PCR assaying rapid formats. The limit of detection of diluted RNAs using MGB probe rRT-PCR assay reached to  $\leq 6$  fg/ul. Besides, high specificity of the former assay was clear. In contrary, the employing of FMDV O specific primer pairs in SYBR Green real-time RT-PCR showed a relative less sensitivity and specificity, particularly one of them displayed a very poor performance. Lastly, the local financial cost of MGB probe is considered the obvious hinder.

Keywords: Foot-and-Mouth disease virus, rRT-PCR, MGB probe

#### **1. Introduction**

Foot-and-mouth disease (FMD) is the most economically significant animal viral disease worldwide affecting cloven-hoofed animals and is caused by Footand-mouth disease virus (FMDV). The virus is a picornavirus (genus *Aphthovirus*, family *Picornaviridae*). The virus has a linear single-stranded RNA genome (*Carrillo et al., 2005*). Seven antigenically distinct forms of the virus are known, called serotypes, but serotype C has not been detected anywhere for many years and may now be extinct. The serotypes have been further divided into topotypes (except for serotype Asia-1 viruses, which comprise a single topotype), genotypes, lineages and sublineages, which are usually restricted to specific geographical regions (*Jamal and Belsham, 2018*).

The rapid and precise detection of FMD virus is a prerequisite. Conventional reverse transcriptase polymerase chain reaction (RT-PCR) (*Amaral-Doel et al.*, 1993; Vangrysperre and De Clercq, 1996; Reid et al., 1998, 2001) and real-time RT-PCR assays (*Callahan et al., 2002; Reid et al., 2002; Moniwa et al., 2007; Tam et al., 2009*) have been developed to complement primary diagnostic techniques for the detection of FMDV. Real-time RT-PCR assays recommended by the World Organization for Animal Health (Office International des Epizooties, OIE) for detection of FMDV incorporate universal primers and fluorescent-labeled probes that recognize conserved regions within the 5-UTR & 3D polymerase (*OIE, 2017*).

The usage of a panel of rRT-PCR assays is imperative, as RNA, viruses are prone to mutation. If one assay is rendered ineffective due to a castrophic

mutation in the primer or probe binding regions or as a result of event of contamination problems, results from other assay will still be valid specially when using assays targeting different areas of the genome (Mckillen et al., 2011). The MGB molecule involved in the detector probe design increase the Tm of the probe. This shortens the probe sequence and enables it to be designed strictly to in variant region. Moreover, the dynamic range of MGB-NFQ probes is larger because of its increased fluorophore quenching, efficiency and resulting low fluorescent background compared to FAM-TAMRA probes (*Moniwa et al., 2007; Kutyavin et al., 2000*).

FMDV serotype O is the most ancient well-identified worldwide type (*Vallée and Carré, 1922*). Also, in Egypt it the classical enzootic and the most prevalent serotype pose many outbreaks (*Aidaros, 2002; Kardjadj, 2018*). Therefore, application and assessing of a new trend for viral detection in Egypt depending on minor groove binder probe (MGB) real-time RT-PCR (rRT-PCR) was applied on local FMDV serotype O, A, and SAT2 with special handling of FMDV O serotype specific primer sets in SYBR Green real-time RT-PCR for the rapid and precise detection of the virus

## 2. Materials and Methods

## 2.1. Viruses

The Egyptian strains of FMDV serotypes O, A and SAT2 were used in this study included O/EGY/2009 iso1 (cell culture grown virus), A/EGY/2009 iso-Cai (clinical isolate) and SAT2/EGY/H1Ghb/2012 (bovine tongue epithelium suspension from Gharbia, Egypt) (*Abu-Elnaga, 2011, EL-Shehawy et al., 2011 and 2014*). They were initially utilized for validation of Minor groove binder probe real-time RT-PCR in detection of different Egyptian FMDV serotypes. Moreover, Other FMD viruses representing FMDV serotypes found in Egypt, previously type discriminated by RT-PCR assay and nucleotide sequence were used as unknown samples in the performance of the probe rRT-PCR assays. These FMD viruses were clinical and culture viruses, O (n=10), A (n=4) and SAT2 (n=2). In context of that, a related vesicular viral disease that cause mucosal lesion with excessive salivation accompanied by lameness in chronic infection was, also, incorporated in the assays. Also, Bovine Viral Diarrhea virus (BVDV, RNA molecule of approximately 12.3 kb) were used to illustrate specificity of the current article assay.

## 2.2. RNA purification and analytic sensitivity

QIAamp® Viral RNA kit (Qiagen, Germany) for RNA extraction was used according to the manufacturer's instruction. Extracted RNAs from three FMD viruses of serotype O were quantitated by UV spectrophotometry and used as; in-house Standard (a candidate culture propagated virus in 2017 with titer ~7 (107 log10) TCID50/ml on BHK), Positive control-1 (semi-purified concentrated culture grown virus in 2012) and Positive control-2 (cell-adapted FMDV isolate in 2009). The standard was imperative to obtain the standard curve. Bulks of extracted RNAs from the standard and positive control 2 were divided in two aliquots for each. From one aliquot of the standard RNA, a seven times, serial 10-fold dilutions in Rnase-free water were performed to obtain the

Standard RNA Dilutions (SRD) to have values that were used to construct a standard curve to calculate unknown sample concentrations.

An archived Stock virus (SV) RNA (Azab et al, 2012), kept for a complete 6-years in an ordinary kitchen fridge and previously assessed by SYBR Green rRT-PCR, was examined in the current article for assessment rRT-PCR assays on RNAs that suffered storage for a long duration. Briefly, 10fold serial dilutions of stock virus (SV) in minimum essential medium (MEM) with Hank's salts in the range of 10-1-10-8 were performed. Each dilution was exposed to RNA isolation procedures to prepare SV RNAs. The previous different RNAs preparation formats viz. SRD and SV RNA was used in analytic sensitivity. Negative controls included: no template control, NTC, which was Rnase free water; Negative control 1 that was RNA from healthy BHK cells; and Negative control 2 that was RNA from non-infected BHK cells showed contamination. For quantification the mass concentration of RNA, two spectrophotometer instruments were utilized, one was the conventional spectrophotometer (Milton Roy 601 Spectronic 335104, USA) and other was the modern spectrophotometer (NanoDrop 2000c Spectrophotometer, Thermo Fisher Scientific, USA).

Repeatability assay of MGB probe rRT-PCR sensitivity was performed after one-month interval on FMDV O RNA standard, its 1<sub>st</sub> dilution (10-1) and positive control 2, which were considered as an old exhausted aliquot RNA after using for one-month. Where the standard and control 2 had been exposed to repetitive freezing and thawing for ten and six times, respectively.

Using the former samples and depending on the current primers flanked the MGB probe, each MGB and SYBR green rRT-PCR were extra re-assessed with the previous old aliquot RNA, but also, new aliquots of the standard, positive control 2 and primers/probe set were additionally involved in the assay. **2.3. Real-time RT-PCR (rRT-PCR)** 

All the extracted RNAs were tested on the real-time PCR system Rotor-Gene Q 2 (Corbett Life Science, a QIAGEN Company, Germany) using either QuantiTect Probe RT-PCR Kit or QuantiTect SYBR Green RT-PCR Kit (Oiagen, Germany). Primer set: PF-5-GTT TTG TTC TTG GTC ACT CCA T-3'; PR-5'-ACG GAG ATC AAC TTC TCC TGT A-3' and a labeled FAM, 5' conjugated minor groove binder (MGB) probe CTC TCC TTT GCA CGC C, 5'- FAM 3'-NFO/MGB, were employed in MGB rRT-PCR investigations with approximately 163 bp target genome fragment amplification (McKillen et al., 2011). The primers were purchased from either Metabion, Germany or Bioneer, Korea; while, the Probe was designed by Applied Biosystems, Life Technologies, Thermo Fisher Scientific, USA. The serotype O specific primer pair (our lab termed PH1/PH2), designed from the 1D and 2AB regions of the viral genome as described previously (Vangrysperre and De Clerca, 1996) to give 402 bp expected band sizes were used in SYBR Green rRT-PCR assay for further MGB comparative assessment. Also, for auxiliary performance evaluation, the previously (Knowles and Samuel, 1994) documented oligos O-1C124 (ARS4)/NK61 of an expected amplicon of approximately 1126 bp were implemented by SYBR rRT-PCR method. For TaqMan MGB rRT-PCR,

cycling conditions were: 1 cycle at 45 oC for 30min, 1 cycle at 95 oC for 15min and 45 cycles at 95 oC for 30s, 51 oC for 30s, and 72 oC for 30s. For the SYBR Green rRT-PCR methods, the optimized reaction contained 5% RNA template. The cycling parameters were as described previously (*Knowles and Samuel, 1994; Vangrysperre and De Clercq, 1996*) and for 45 cycles. At least one of the positive controls and negative controls was involved in every assay.

## 3. Results

MGB rRT-PCR assay were valid to detect the specific target genome fragment of different Egyptian FMDV serotypes strains. Besides, various negative controls and the tested BVDV produced neither threshold cycle (CT) values nor the expected amplification sizes in agar gel electrophoresis. Using MGB amplification, RNA sensitive detection of the standard and all its dilution extended to the theoretical mass quantification of six hundred attograms (ag) RNA per microliter ( $\mu$ I).Also, sensitivity were attained in examining the archived stock virus (SV) RNA, MGB fluorogenic signals were attained in dilutions from 10-1 to 10-4 and 10-6. On the other hand, MGB probe and primers failed to hybridize to the respective templates in dilutions 10-5, 10-7 and 10-8 (**Fig. 1 and 2**).

SYBR green rRT-PCR method using primer pair for FMDV serotype O (**Fig. 3**), showed specific sensitivity reached to sixty femtogram RNA per microliter utilizing PH1/PH2 primer set. Furthermore, quantification of six hundred picogram (might be barely reach 60 pg in repeatability) of RNA per microliter were obtained when employing alternative oligos for FMDV O, 1C124 (ARS4)/NK61 (**Fig. 4**).

Likewise, Repeatability of MGB rRT-PCR assay revealed variability in the sensitivity between aliquots and replicates with a highlighted effect of RNA degradation and an approximately negligible impact of the primers /probe hybridization regression. CT values variability using MGB probe between the replicates for each sample in comparison with the mean was  $\pm 3.5$  with values difference reached 6.5, 5.7, and 2.9 for Positive Control 2, Standard and Standard 10-1, respectively (**Fig. 5 and 6**) and Table **1.** Accordingly, it was suggested to be the cut off  $\leq$  45 cycles for MGB probe due to its high sensitive and specific detection.

Using the primer set that flanked the MGB probe in either Probe or SYBR rRT-PCR methods, successively at the same day, clearly showed the performance of the standard and its dilution were promising in Probe in comparison to SYBR assays, where the signals were linear in the exponential phase. The standard curve efficiency for the MGB probe was 0.89, while for the SYBR PH1/PH2 primers was 0.79. Using Taqman MGB Probe, the standard virus and its serially 10-fold dilutions showed a 2.8-4.4 increment increasing of CT values between undiluted virus until its 10-6 dilution (undiluted & 6 dilution. Furthermore, using SYBR Green PH1/PH2 oligos protocol, the standard virus and its serially 10-fold dilutions gave CT values of 3.3-4.6 differences between undiluted virus until its 10-6 dilution (undiluted & 6

dilution series). Melt curve peak (Tm) showed that specific amplification giving the expected peak, affirmed by yielding the anticipated fragment size on agarose based electrophoresis. The negative samples controls either did not exhibit the anticipated specific former Tm of the primers or showed the expected peak, but they all had higher CT values above the last positive standard dilation

#### 4. Discussion

Shorter, more specific minor groove binders (MGBs) probes are dsDNAbinding agents attached to the 3' end of TaqMan probes to increase the Tm value (by stabilization of hybridization) and to design shorter probes. Shorter probes make it easier to use short conserved or unique sequences for hybridization. MGBs also reduce background fluorescence and increase dynamic range due to increased efficiency of reporter quenching due to shorter distances between the reporter and quencher and the use of non-fluorescent (dark) quenchers (NFQ) at the 3' end instead of fluorescence dyes like TAMRA. MGB probes have more sequence specificity for better mismatch recognition.

The format of standard and its dilutions (SRD) in the current article was selected according to what comprehend from the thermal cycler manufacture's recommendation, which mentioned that the DNA used in the standard curve should be derived from similar DNA in the samples being measured. It was recommended that the concentration of at least one DNA sample be determined using ultraviolet spectrophotometry and that this sample be used as the standard. The minimum number of standards used should be three (with replicates). Importantly, DNA standards used in fluorescence detection are only linear within the range of 100 nano-grams per micro-litre to 1 nano-gram per micro-litre. That is, within this range, if the concentration of DNA is halved, so is the fluorescent reading. The confidence intervals for any concentration outside this range are very broad due to non-linearity in the chemistry. In addition, differences have been observed in the measurement of various forms of DNA. For example, genomic DNA compared with plasmid DNA. Therefore, it is recommended that only alike DNA are measured together, and the use of plasmid DNA as a standard be avoided when measuring genomic DNA.

The oligonucleotide probe with a 5' conjugated minor groove binder (MGB) ligand as a reporter in real-time PCR. The hybridization of the probe triggered fluorescent. MGB probe rRT-PCR was the best specificity and sensitivity than the other two primer pairs used in SYBR Green RT-PCR protocols. Relevant amplification of the homologous templates were implemented, whereas, the heterologous templates were mismatched. Consequently, the specificity of this fluorogenic probe was very satisfactory for FMDV investigation and quantification. The fluorogenic MGB probes were more specific for single base mismatches and fluorescence quenching was more efficient, giving increased sensitivity (*Kutyavin et al, 2000*). Result revealed lower CT values, in addition to, higher detection specificity and sensitivity when using FMDV O specific primers that produce smaller amplification size, in comparison to, O specific

oligos amplifying larger fragment. In real-time PCR with TaqMan probes, the amplicon size directly influenced detection: the larger the amplicons, the later the detection. Earlier detection and a higher fluorescence level (plateau phase) were generally observed for shorter amplicons (*Debode et al., 2017*). On the other side in SYBR Green methods, melting curve analysis was considered as a tool to verify the specificity of the amplified product, although it is a common indicator used in fluorescence rather than fluorophore-based RT-PCR assays. Besides, agar gel electrophoresis supports the amplicon specificity. In contrary to MGB probe, the serotype specific oligos, 1C124 (ARS4)/NK61, trial in SYBR rRT-PCR format gave the poorest analytical specificity and sensitivity.

Analytical sensitivity was a trial to detect variable genome of FMDV of different serotype to overcome the possibility of false negative result due to serotype unspecificity. In the absence of a target molecule, the MGB probe does not fluoresce, as there is sufficient interaction between the reporter fluorophore and the quencher to prevent a fluorescent signal. Hybridisation to a complementary target molecule triggers an increase in fluorescence due to the separation of the fluorophore and quencher (*Mckillen et al., 2011*). Probe with minor grove binder (MGB) form stable, higher melting temperature interactions with their target sequences. The positive result of rRT-PCR in this article are mainly CT values. Where there is general correlation between them and quantity of input nucleic acid. The more target template is present in the reaction, the fewer cycles it require to reach logarithmic growth and end point of RT-PCR (i.e. lower CT values) (*Moniwa et al., 2007*).

After one month of  $1_{st}$  assessment of the performance and the analytic sensitivity of MGB rRT-PCR, an anticipated detection limit decreased by 10-fold, with 2.6-5.5 increment difference in CT values between undiluted virus until its 10-6 dilution, where higher increment values in lower dilutions and lower increment values in higher dilution. The operator for faulty cost saving did not carry out re-assessment of dilution 10-7. However, result was satisfactory by ending the re-assessment by dilution 10-6 and not exceed to dilution 10-7, because the signal curve of dilution 10-6 was at the border of the threshold (CT = 43) that predict if dilution 10-7 was done, the CT value would be weak positive or negative.

Six-years before the current assay, SYBR Green rRT-PCR had detection limit for the stored Stock virus (SV) RNA that was extended to 10-7 dilutions using 0.2 RNA template/rxn volume (Azab et al., 2012). In context, the current MGB probe was still cable to detect RNA in the archived RNA across approximately 6-log range of input template with 1 log10 regression, taking in consideration that the RNA template input in this paper was 4x lesser. Also, the detection signals produced as a result of SYBR Green DNA incorporation were generally more earlier as revealed in repeatability assay when MGB probe complementary primers were used in SYBR Green investigation. This result was satisfactory when investigation a long period storage of RNA template.

The nucleic acid amplification detection on the real-time PCR platform was verified by agarose gel electrophoresis that revealed the expected positive band. Serial dilution of the virus RNA control could be used as one of the viable reference for relative FMDV quantification. The virus RNA control sample was of cultured derived virus, not a wild virus to minimize the possibility of contamination by non-specific fragment. Furthermore, in future we hope to use in-vitro synthetic FMD RNA fragment of the primer/probe target sequence as a positive amplification control.

The drawback of the current MGB probe (FAM dye–labelled, with NFQ), in comparison to the non-MGB assay (FAM dye–labeled, with BHQ) in the poor developing country is the relative comparative higher cost of MGB probe. That might reached to 1.7x the price of non-MGB probe that will be translated to thousands of EGP (or hundred of USD), in consequence of that, rising the finance of the quantity detection assay of the unknown samples.

Finally, MGB RT-PCR assay provided a rapid, sensitive, specific and less labor for detection of FMDV with subsequent early planning for a control strategy in case of an outbreak with liberating FMDV free animals from quarantine measure

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## **Conflict of Interest**

The authors declare that they have no conflict of interest.

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Conditions 1 Time; Probe or SYBR	1 <sub>st</sub> day probe same run		1	2nd day SYBR same run				
Conditions 2 rxn mixture content	old primers /probe aliquot		ers uot	new primers aliquot				
Conditions 3 rxn mixture content	new RNA aliquo t							
Replicate no	2							

Table 1: CT values of three samples exposed to different conditions

Positive Control 2	27.7					
Standard	25.2					
Standard, 10-1	ND					
NTC	-					

.ND=not done, Reading was CT values recorded in various conditions



Fig. 1. Amplification sensitivity of MGB probe rRT-PCR for FMDV O RNA prequantified, 6 ng/ $\mu$ l, and serially 10-fold diluted until theoretically reach 6 ag/ $\mu$ l. The

former virus was used as standard. Nomenclature on the real-time figure is representing curves either by intersecting the 1st digit (i.e. 6) in the quantity number of the RNA masses or by having an annotation with a black shadow pointed to it. The threshold is the horizontal red line intersecting the curves. (A) Quantitation data of the standard and its dilutions. Two positive controls, two negative controls, no template control (NTC) and non-specific virus (BVD) were involved in the assay run. Positive CT values were curves peaks above the threshold, while negative values were peaks at the borders or below the threshold (B) Agar-based electrophoresis of the amplified products by the MGB probe. M: 100 bp ladder. Expected fragment size were approximately 163 bp of FMDV 3D gene. Lanes 1-8: The standard dilutions from 6 ng/µl to 6 ag/µl, Lanes 9-10: negative control 1 & 2, Lanes 11: BVD, Lane 12: NTC, Lanes 13-17: Unknown samples, that were skipped in Quantitation data figure to not hide the illustrated standard curves. Positive controls were not shown.



**Fig. 2.** (A) Detection of an archived FMDV stock virus (SV) RNA by MGB probe rRT-PCR. SV dilutions 10-1-10-8, except 10-3, showed various CT values with amplification cycles. Two positive and negative controls, NTC, non-specific virus (BVD) and unknown samples were involved in the test. (B) Detection of stock virus (SV) RNA with anticipated 163 bp. M: 100 bp ladder. Lanes: 1-7 SV dilutions 10-1-10-8, except the not done dilution 10-3. Lanes 8 & 13 were positive control 1& 2. Other lanes were unknown samples, Lanes 16-17: showed a very faint bands of an archived unknown samples, which were amplified by another TaqMan probe of an expected size of 107 bp and were used as a control of electrophoresis





**Fig. 3.** Implementation of SYBR Green rRT-PCR using oligos targeting FMDV O 1D gene of an expected 402 bp for minimum detecting of FMDV O RNA pre-quantified as 6 ng/ $\mu$ l, serially 10-fold diluted and theoretically extended to 6 ag/ $\mu$ l. The former virus was used as standard. (**A**) Graphical representation of real time one-step RT-PCR data of the standard and its dilutions. Negative control, NTC and non-specific virus, BVD, were involved in the assay run. Positive CT values were curves peaks above the threshold, while negative values were peaks at the borders or below the threshold (**B**) Electrophoresis on agarose gel of the rRT-PCR assay samples using SYBR Green. M: 100 bp ladder. Positive bands were approximately 402 bp of FMDV O 1D gene. Lanes 1-8: The standard dilutions from 6 ng/ $\mu$ l to 6 ag/ $\mu$ l, Lane 9: negative control 2, Lane 10: BVD, Lane 11: NTC, Insets show melting curve analysis.



**Fig. 4.** SYBR Green rRT-PCR amplification of an expected 1301 bp embarrassing FMDV O 1D gene of for assessment the detection limit of FMDV O RNA prequantified as 6 ng/µl, serially 10-fold diluted and theoretically extended to 6 ag/µl. The former virus was used as standard. (**A**) Graphical representation of real time onestep RT-PCR data of the standard and its dilutions. Two positive controls, two negative controls, no template control (NTC) and non-specific virus (BVD) were involved in the assay run.Positive CT values were curves peaks above the threshold, while negative values were peaks at the borders or below the threshold (**B**) Electrophoresis on agarose gel of the rRT-PCR assay samples using SYBR Green. M: 100 bp ladder. Positive bands were approximately 1301 bp of FMDV O 1D gene. Lanes 1-8: The standard dilutions from 6 ng/µl to 6 ag/µl, Lane 9-10: negative control 1 &2, Lane 11: NTC, Lane 12: BVD, Lanes 13-16: unknown samples, Melting curve analysis (insets) revealed amplification specificity.





**Fig. 5.** Re-assessment (1-month later from previous evaluation) of the sensitivity of MGB probe rRT-PCR for FMDV O RNA standard (6 ng/ $\mu$ l, and serially 10-fold diluted). (A) Quantitation data of the standard and its dilutions. Unknown samples were involved in the assay run. (B) Agar-based electrophoresis of the amplified products by the MGB probe. M: 100 bp ladder. Expected fragment size were approximately 163 bp of FMDV 3D gene. Lanes 1-7: The standard dilutions from 6 ng/ $\mu$ l to 6 fg/ $\mu$ l. Unknown samples were not shown.



**Fig. 6.** Re-testing, 1-month later from previous current article evaluation, of a standard & its 1<sub>st</sub> RNA dilution for investigate the efficiency of a MGB probe rRT-PCR under variable factors. Positive control 2 using the primers either with MGB probe rRT-PCR **Fig 6A (i)-(iii), Fig 6B (i) or with** SYBR Green rRT-PCR **Fig 6A (iv)-(v), Fig 6B (iii)** depending on an old & new RNA aliquot of the standard and positive control 2, also an old & new aliquots of primers & probe. (A) Real-time detected signals curve (B) Agar gel based electrophoresis

الملخص العربي

اختبار النسخ العكسي و تفاعل البلمرة المتسلسل اللحظي مستخدما مجس رابِط بالأخدود الأصغر لاكتشاف فيروس مرض الحمى القلاعية في مصر هاني إبراهيم أبو النجا معد بحوث الأمصال واللقاحات البيطرية. العباسية القاهرة ص.ب:131

إن مجس رابط بالأخدود الأصغر MGBs، يعد الأقصر والأكثر تحديدًا و الذي يمكن تصميمه بدقة في مناطق جينية مختلفة. تم تطبيق وتقييم اتجاه جديد للكشف عن فيروس مرض الحمى القلاعية في مصر اعتمادًا على تقنية اختبار النسخ العكسي و تفاعل البلمرة المتسلسل اللطي MGB على الأنواع المحلية المختلفة (O, A, and SAT2) للفيروس مع التركيز على الفيروس نوع O و الكشف عنه مستخدما تفاعل البلمرة و SYBR green و فد بلغ حد الكشف عن الحمض النووي باستخدام اختبار MGB إلى اقل من 6 فيمتوجر ام/ميكرليتر. إلى جانب ذلك، كانت خصوصية التجربة السابقة واضحة .على العكس من ذلك ، فإن توظيف أزواج بوادئ للحمض النووي لنوع O في تفاعل البلمرة اللحظي، أظهر حساسية و خصوصية التعتبر المعوق الواضح.