Capture of dengue viruses using antibody-integrated graphite-encapsulated magnetic beads produced using gas plasma technology

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Abstract. Despite significant advances in medicine, global health is threatened by emerging infectious diseases caused by a number of viruses. Dengue virus (DENV) is a mosquito-borne virus, which can be transmitted to humans via mosquito vectors. Previously, the Ministry of Health, Labour and Welfare in Japan reported the country’s first domestically acquired case of dengue fever for almost 70 years. To address this issue, it is important to develop novel technologies for the sensitive detection of DENV. The present study reported on the development of plasma-functionalized, graphite-encapsulated magnetic nanoparticles (GrMNPs) conjugated with anti-DENV antibody for DENV capture. Radiofrequency wave-excited inductively-coupled Ar and ammonia gas plasmas were used to introduce amino groups onto the surface of the GrMNPs. The GrMNPs were then conjugated with an antibody against DENV, and the antibody-integrated magnetic beads were assessed for their ability to capture DENV. Beads incubated in a cell culture medium of DENV-infected mosquito cells were separated from the supernatant by applying a magnetic field and were then washed. The adsorption of DENV serotypes 1-4 onto the beads was confirmed using reverse transcription-polymerase chain reaction, which detected the presence of DENV genomic RNA on the GrMNPs. The methodology described in the present study, which employed the plasma-functionalization of GrMNPs to enable antibody-integration, represents a significant improvement in the detection of DENV.

Introduction

Dengue virus (DENV), a plus-strand RNA virus with an enveloped icosahedral nucleocapsid, can be transmitted through mosquito vectors, and is the causative agent of dengue fever, dengue hemorrhagic fever and dengue shock syndrome (1). Dengue hemorrhagic fever and dengue shock syndrome are potentially life threatening, and the risk of developing these diseases is correlated with infection by one of the four DENV serotypes (DENV1-4) and the carrying of antibodies to another DENV serotype from a previous infection (2). Currently, no specific treatment or vaccine for DENV is available (3).

There has been a significant increase in the number of reports of DENV-associated infections and DENV-associated mortality (1). Notably, over the last 50 years, the incidence of DENV-associated infections has increased by 30-fold, and the World Health Organization estimate that there are currently 50,000,000 cases per annum worldwide (3). Infections due to DENV are now of serious concern worldwide, particularly in tropical areas, including Southeast Asia (4). DENV was first isolated by Kimura and Hotta from blood samples obtained during the 1943 dengue epidemic, which was predominantly confined to the Japanese port cities of Nagasaki, Kobe and Osaka (5). On the 28th August 2014, the Ministry of Health, Labour and Welfare in Japan reported the country’s first domestically acquired case of dengue fever for almost 70 years (6). Therefore, the implementation of procedures to monitor for any further potential outbreak of dengue fever in the country are required.

Consequently, the development of effective methods for the surveillance of DENV are urgently required. One approach to achieve sensitive detection of DENV is to establish a method to concentrate the viral particles. Several approaches...
to concentrate virus have been suggested, including ultracentrifugation and polyethylene glycol (PEG)-mediated precipitation. Although these methods are applicable to a number of viruses, they have significant practical limitations. Specifically, ultracentrifugation is time-consuming and can increase the false-positive rate when combined with polymerase chain reaction (PCR) analysis (7). Although PEG-mediated precipitation is simple and easy to perform, PEG interferes with the subsequent PCR procedure (8,9).

An alternative to these conventional methods is the use of magnetic beads coated with molecules, which efficiently bind to the virus, which allows the capture and concentration of the viral particles by applying a magnetic field. A potential approach to the capture of a target virus is to use magnetic beads coated with an antibody specific to the particular virus of interest.

Magnetic nanoparticles (MNPs), including iron, nickel or cobalt, have been widely investigated for biomedical and environmental applications due to their high specific surface area and the ease of magnetic collection of target materials adsorbed by the MNPs (10,11). However, a significant problem with these beads is their inherent chemical instability, which can limit their application in the field of biological and environmental science (12). To overcome this limitation, the MNPs are typically encapsulated with a protective shell of graphite, silica or polymer (12).

Graphite-encapsulated MNPs (GrMNPs) are usually hydrophobic, which is a limitation for several biomedical applications. However, appropriate surface modification can improve the properties of the GrMNPs, allowing them to efficiently recognize and bind to molecular targets, including antibodies, antigens and receptors (13). Amino group functionalization, which is a desirable functionality for graphite, improves the reactivity and hydrophilic nature of the GrMNPs (12). A promising method for the amino functionalization of GrMNPs is to use inductively coupled radiofrequency (RF) plasma, which is environmentally friendly and requires a short duration for the reaction to reach completion. Using this approach, it is possible to introduce amino groups effectively (14). Furthermore, the degree of surface derivatization with amino groups of GrMNPs can be optimized by altering the plasma discharge conditions (15). With these amino-modified GrMNPs, the efficient surface immobilization of sugar chains has been demonstrated, including dextran, as well as antibodies against several pathogens, including anti-influenza virus and anti-Salmonella antibodies (12,16,17).

On the basis of this background, the present study was performed to expand on previous results examining the influenza virus to investigate DENV via the immobilization of anti-DENV antibody onto the functionalized surface of GrMNPs. The modified GrMNPs were then assessed for their ability to capture DENVs, and the concentrated virus was then detected in combination with a PCR-based amplification procedure.

Materials and methods

Plasma-functionalized GrMNPs and production of antibody-integrated magnetic beads. The graphite-encapsulated iron compound nanoparticles were prepared using an arc discharge method by applying a 150-200 A direct current at ~20 V between an anode and cathode, as described previously (15). A graphite electrode, molded using graphibond-551R with Fe₃O₄ powder, was used as the anode. On the opposite side, a graphite rod (50 mm×10 mm; 99.9%) was used as the cathode. The resulted graphite-encapsulated iron compound nanoparticles were then exposed to plasma, which was produced using an RF power supply (18,19) in an atmosphere containing ammonia at 13.56 MHz and 80 W via a matching network (18,19). Initial pretreatment was performed for 10 min using Ar plasma, followed by 2 min of ammonia plasma post-treatment for amino group introduction. During the experiments, the gas pressure was maintained at 50 Pa. The amino groups on the surface of the magnetic beads were then further labelled with 0.3 μM of the coupling agent, N-succinimidyl 3-(2-pyridyldithio) propionate (SPDP; Dojindo Laboratories, Kumamoto, Japan) at pH 7-8. A human monoclonal antibody (clone no. D23-1G7C2) recognizing the first domain II fusion region of the DENV envelope glycoprotein (E) (20) was reduced using diithiothreitol (DTT), resulting in breakage of the S-S bonds and generation of S-H groups. The D23-1G7C2 antibody was produced from hybridomas using peripheral blood mononuclear cells from patients in the acute phase of dengue fever 5 days following the onset of illness, and exhibits neutralizing activity against DENV1-4 (20). The S-H groups on the antibody were then reacted with the SPDP-NH₂-magnetic beads, resulting in covalent crosslinking of the antibody onto the surface of the beads. The resulting magnetic beads were termed antibody-integrated magnetic beads (Fig. 1).

Cell culture and virus. A C6/36 cell culture (American Type Culture Collection, Manassas, VA, USA), derived from Aedes albopictus, was maintained in Leibovitz L15 medium (Thermo Fisher Scientific, Inc., Waltham, MA, USA) containing 0.3% tryptose phosphate broth (TPB) and 10% fetal calf serum (FCS; Wako Pure Chemical Industries, Ltd., Osaka, Japan). The laboratory DENV strains (21), DENV1 (Mochizuki strain), DENV2 (16681 strain), DENV3 (Type Culture Collection, Manassas, VA, USA), derived from Salmonella enterica serovar Typhi (S. Typhi) by the procedure of a previous report (12). DENV3 (80-2 strain) and DENV4 (H241 strain), were used to infect the C6/36 cell cultures. The C6/36 cells were cultured to ~80% confluence, then infected with the DENVs at a multiplicity of infection of 0.1 in Leibovitz L15 medium containing 0.3% TPB and 2% FCS, and were incubated for 3 days at 28°C. The medium was then collected and used for viral capture experiments.

DENV capture. The capture of DENV1-4 was performed as follows. Briefly, 10 μl of the magnetic beads were washed twice with phosphate-buffered saline (PBS). A 10 μl sample of medium from uninfected (Mock) or DENV-infected cell cultures were added to the washed beads with 1 ml PBS, and the tube was incubated for 15 min at room temperature. The tubes containing the mixtures were then set in a magnetic field for 5 min using an Adem-Mag SV magnetic device (Ademtech, Pessac, France). Following magnetic separation, the supernatant was removed, and the beads were washed three times with PBS and resuspended in 10 μl PBS. This procedure produced two fractions: Bead fraction (BD) and supernatant fraction
Ammonia group detection using fluorescamine. Flurescamine reacts with amino groups and forms blue-green fluorescent derivatives. In the present study, fluorescence intensity increased in a dose-dependent manner with bovine serum albumin (BSA) concentrations following incubation with fluorescamine (Wako Pure Chemical Industries, Ltd.) at an excitation of 400 nm and emission of 490 nm. The amino groups on the GrMNPs treated with ammonia plasma were measured using a fluorometric method with fluorescamine, in accordance with a previous study with modification (23). Briefly, 5 µl of the plasma-treated nanoparticles (2 mg/ml) were solubilized with 100 µl 50 mM borate buffer, following which 4 µl 0.075% fluorescamine in acetonitrile was added to each sample. The fluorescence, at an emission of 490 nm and excitation of 400 nm, was measured using a SpectraMax M2e HK fluorometer (Molecular Devices, Sunnyvale, CA, USA).

Amino group detection using 2,4,6-trinitrobenzenesulfonylic acid (TNBS). TNBS reacts with amino groups and shows an increase in absorbance at ~350 nm (24). When 10 mg/ml BSA was reacted with TNBS (Wako Pure Chemical Industries, Ltd.), the maximal peak of the spectra was 345 nm. In addition, a dose-dependent increase of the corresponding peak in BSA was observed following the TNBS reaction at 345 nm. The amino groups on the GrMNPs treated with ammonia plasma were measured using a colorimetric method using TNBS in accordance with a previous study with modification (24). Briefly, 20 µl of the plasma-treated GrMNPs (2 mg/ml) in 50 mM borate buffer was incubated for 16 h at 4°C following the addition of 0.5% TNBS. Subsequently, 250 µl of 5% sodium dodecyl sulfate and 2 N HCl were added, and the absorbance at 345 nm was measured using a spectrophotometer (UVmini-1240; Shimadzu Corporation, Kyoto, Japan).

RT-PCR for DENV. Viral genomic RNA was extracted from the samples prior to and following magnetic separation using a QIAamp Viral RNA mini kit (Qiagen). The RNA was reverse transcribed using a PrimeScriptII first strand cDNA synthesis kit (Takara Bio, Inc.) with random primers. Following incubation at 65°C for 5 min, the viral RNA was reverse transcribed at 42°C for 60 min. The resultant cDNA (2 µl) was amplified by PCR, in a reaction mixture (20 µl) containing primers (0.2 µl each, 100 pmol/µl), MgCl2 (1.6 µl, 25 mM), dNTP mixture (1.6 µl, 2.5 mM each), Ex Taq (0.1 µl, 5 U/µl; Takara Bio, Inc.) and 10X Ex Taq buffer (2 µl) under conditions of 35 cycles at 94°C for 30 sec, 55°C for 30 sec and 72°C for 1 min. The primers used were as follows: D1, forward 5'-ctctaagctgaaacggaggaaacg-3' and D2, reverse 5'-tgactaaagctgctgcaggttcggttt-3' (22). The amplified DNA fragments were purified and analyzed by DNA sequencing on an ABI PRISM3100 genetic analyzer (Applied Biosystems; Thermo Fisher Scientific, Inc.) to verify the identity of the amplified product. Then, the product sequences were compared with the sequences in the Genbank database (ncbi.nlm.nih.gov/genbank/).

The RT-PCR analysis was performed using a QIAamp Viral RNA mini kit (Qiagen, Hilden, Germany), according to the manufacturer's protocol. The RT reactions were performed using a PrimeScriptII First Strand cDNA synthesis kit (Takara Bio, Inc., Otsu, Japan) with random primers. Following incubation at 65°C for 5 min, the viral RNA was reverse transcribed at 42°C for 60 min. The resultant cDNA (2 µl) was amplified by PCR, in a reaction mixture (20 µl) containing primers (0.2 µl each, 100 pmol/µl), MgCl2 (1.6 µl, 25 mM), dNTP mixture (1.6 µl, 2.5 mM each), Ex Taq (0.1 µl, 5 U/µl; Takara Bio, Inc.) and 10X Ex Taq buffer (2 µl) under conditions of 35 cycles at 94°C for 30 sec, 55°C for 30 sec and 72°C for 1 min. The primers used were as follows: D1, forward 5'-ctctaagctgaaacggaggaaacg-3' and D2, reverse 5'-tgactaaagctgctgcaggttcggttt-3' (22). The amplified DNA fragments were purified and analyzed by DNA sequencing on an ABI PRISM3100 genetic analyzer (Applied Biosystems; Thermo Fisher Scientific, Inc.) to verify the identity of the amplified product. Then, the product sequences were compared with the sequences in the Genbank database (ncbi.nlm.nih.gov/genbank/).

Statistical analysis. Statistical analyses were performed using Prism 4 software (GraphPad Software, Inc., San Diego, CA, USA). The TNBS and fluorescamine data were subjected to a non-repeated measures analysis of variance followed by the Bonferroni correction test. The PCR data were subjected to Student's t-test (unpaired). P<0.05 is considered to indicate a statistically significantly difference.

Figure 1. Schematic representation of the ammonia plasma-treated GrMNPs and their binding to anti-DENV antibody, resulting in production of antibody-integrated magnetic beads. The surfaces of the GrMNPs were reacted with ammonia plasma, produced using a radiofrequency power supply, resulting in the introduction of amino groups (NH2-beads) at pH 7-8. Anti-DENV antibody (D23-1G7C2) was reduced using dithiothreitol, resulting in the breakage of S-S-bonds and the generation of S-H groups. The S-H group of the antibody was then further reacted with the SPDP-NH2-magnetic beads. The resultant magnetic beads were termed antibody-integrated magnetic beads. GrMNPs, graphite-encapsulated magnetic nanoparticles; DENV, dengue virus; SPDP, N-succinimidyl 3-(2-pyridylidithio) propionate.
Results

In order to analyze the hydrophilicity of the ammonia plasma-treated GrMNPs, the resultant NH₂-beads were incubated with various concentrations of SPDP (Fig. 2). The results showed that the beads aggregated following incubation with 30 µM SPDP, whereas the beads incubated with 0.03, 0.3 and 3 µM SPDP remained in suspension. These observations suggested that the ammonia plasma-treated GrMNPs possessed hydrophilic properties, possibly due to the introduction of amino groups to the bead surface.

To further confirm the presence of amino groups on the surface of the beads, the ammonia plasma-treated GrMNPs were incubated with TNBS or fluorescamine, which react with an amine functionality. The absorbance at 345 nm was then measured. As a positive control, samples of 0-10 mg/ml BSA were incubated with TNBS for 5 min and their absorbance measured. The absorbance of the ammonia plasma-treated GrMNPs + TNBS prior to and following centrifugation was significantly higher than that of the buffer + TNBS group, indicating TNBS had reacted with the amino groups on the plasma-treated GrMNPs. BSA measurements were obtained as positive controls (n=1). *P<0.05, compared with buffer + TNBS. Values are presented as the mean ± standard error. GrMNPs, graphite encapsulated magnetic beads; TNBS, 2,4,6-trinitrobenzenesulfonic acid; BSA, bovine serum albumin.

Table I. Confirmation of amino groups on the surface of ammonia plasma-treated GrMNPs by reaction with TNBS.

| Treatment | Absorbance at 345 nm |
|-----------|----------------------|
| Buffer + TNBS | 0.2973±0.0002       |
| Ammonia-plasma-treated GrMNPs + TNBS | 0.8153±0.004* |
| Supernatant of ammonia-plasma-treated GrMNPs + TNBS (post-centrifugation) | 0.6833±0.003* |
| BSA (0 mg/ml) | 0.295               |
| BSA (0.1 mg/ml) | 0.327              |
| BSA (1 mg/ml) | 0.499               |
| BSA (10 mg/ml) | 1.745               |

TNBS was used to detect amino groups on ammonia plasma-treated GrMNPs. TNBS was incubated with either the ammonia plasma-treated GrMNPs in borate buffer prior to centrifugation, the supernatant following centrifugation, or with borate buffer alone. The absorbance at 345 nm was then measured. As a positive control, samples of 0-10 mg/ml BSA were incubated with TNBS for 5 min and their absorbance measured. The absorbance of the ammonia plasma-treated GrMNPs + TNBS prior to and following centrifugation was significantly higher than that of the buffer + TNBS group, indicating TNBS had reacted with the amino groups on the plasma-treated GrMNPs. BSA measurements were obtained as positive controls (n=1). *P<0.05, compared with buffer + TNBS. Values are presented as the mean ± standard error. GrMNPs, graphite encapsulated magnetic beads; TNBS, 2,4,6-trinitrobenzenesulfonic acid; BSA, bovine serum albumin.

Table II. Confirmation of amino groups on the surface of ammonia plasma-treated GrMNPs by reaction with fluorescamine.

| Treatment | Fluorescence (excitation 400 nm; emission 490 nm) |
|-----------|-----------------------------------------------|
| Ammonia-plasma-treated GrMNPs | 112.49±7.71* |
| Antibody-integrated magnetic beads | 101.00±7.29* |
| BSA (0 mg/ml) | 84.23±2.55 |
| BSA (0.1 mg/ml) | 178.28±15.88* |
| BSA (1 mg/ml) | 862.58±5.70* |

Fluorescamine was incubated with either ammonia plasma-treated GrMNPs in borate buffer or borate buffer alone, and fluorescence was measured. As a positive control, samples of 0-1 mg/ml BSA were incubated with fluorescamine for 5 min and their fluorescence signals were measured. The results confirmed the amino groups reacted with fluorescamine. The fluorescence signals of the ammonia plasma-treated GrMNPs and anti-dengue virus antibody-integrated magnetic beads were significantly higher, compared with the borate buffer alone. These observations indicated that fluorescamine had reacted with the amino groups present on the surface of the plasma-treated GrMNPs. *P<0.05, compared with 0 mg/ml BSA. Values are presented as the mean ± standard error. GrMNPs, graphite encapsulated magnetic beads; BSA, bovine serum albumin.

Figure 2. Aggregation and suspension of ammonia plasma-treated GrMNPs following incubation with SPDP. A 0.2 mg/ml suspension of amino group-functionalized magnetic beads (NH₂-beads) was incubated with 0.03, 0.3, 3 or 30 µM SPDP for 30 min at room temperature. Aggregation of the beads was observed following incubation with 30 µM SPDP, whereas the beads were in suspension in the 0.03, 0.3 and 3 µM SPDP solutions. These observations indicated the appropriate concentration of SPDP for the suspension of beads and suggested they exhibited hydrophilic properties due the presence of amino groups at their surface. GrMNPs, graphite-encapsulated magnetic nanoparticles; SPDP, N-succinimidyl 3-(2-pyridyldithio) propionate.
following incubation of BSA with the respective reagents. Subsequently, the ammonia plasma-treated GrMNPS were exposed to either TNBS or fluorescamine. The ammonia plasma-treated GrMNPs exhibited increased absorbance at 345 nm and increased fluorescent intensity (excitation 400 nm; emission 490 nm) following incubation with TNBS and fluorescamine, respectively. Following incubation with TNBS, the samples were centrifuged for 5 min at 20,000 x g and room temperature, and the resultant supernatant also showed increased absorbance at 345 nm, indicating that the increased absorbance was not due to turbidity. Taken together, these results suggested that the ammonia plasma-treated GrMNPs possessed surface-exposed amino groups. Subsequently, the present study examined the functionality of the resultant antibody-integrated magnetic beads by incubating them with a suspension of DENV prior to separation of the beads by applying a magnetic field to confirm capture of the viral particles (Fig. 3). Specifically, media from C6/36 mosquito cells infected with DENV1-4 were diluted with PBS and mixed with the magnetic beads. The mixture was then magnetically separated into the BD and supernatant SP fractions (Fig. 4). Medium from mock-infected cells was used to prepare the
control fractions. Finally, the fractions were analyzed using RT-PCR to determine the extent of DENV capture by the beads.

Using primers specific for a 511 bp section of DENV RNA, RT-PCR was used to amplify a product of the expected size in the BD fraction, but not in the SP fraction in DENV1-4 (Fig. 4). Two independent DNA sequence analyses were performed to confirm the identity of the 511 bp band. In each case, within the sequenced region, the following results were obtained using the Genbank database: DENV1 (Mochizuki strain) exhibited 98 and 99% sequence identity to Genbank accession no. AB074760; DENV2 (I6681 strain) exhibited 98 and 99% sequence identity to Genbank accession no. U87411; DENV3 (80-2 strain) exhibited 95 and 95% sequence identity to Genbank accession no. AF317645; and DENV4 (H241 strain) exhibited 99 and 99% sequence identity to Genbank accession no. AY947539.

Subsequently, viral RNA was extracted, transcribed and subjected to RT-PCR using specific forward and reverse primers for DENV in order to investigate the quantity of DENV genomic RNA prior to and following magnetic separation (Fig. 5). The percentage of genomic RNA in DENV1 recovered following magnetic separation by the antibody-integrated magnetic beads was 310.7±46.0%, compared with the control. Of note, this sample was the same as the BD fraction. By contrast, the percentage in the sample prior to incubation with the beads was 97.3±11.2%, compared with the control. These findings suggested that the antibody-integrated magnetic beads concentrated the DENV1 by 3.2-fold.

The above results confirmed that the BD fraction contained the corresponding DENV genomic RNAs, confirming that the beads had successfully captured and concentrated the DENV particles.

Discussion

Our previous studies showed that influenza virus, an enveloped RNA virus, and Salmonella enterica, a pathogenic bacterium associated with food poisoning can be efficiently captured using amino modified GrMNs, generated by an ammonia gas plasma mediated strategy, coated with the corresponding antibody (16,17). Therefore, the generic strategy of using ammonia gas plasma to introduce amino groups into GrMNP, and then covering the beads with antibodies directed against a specific bacteria, virus or other pathogen, facilitates their efficient capture from liquid samples. The targeted pathogen is then pre-concentrated using immunomagnetic separation, enabling their identification using a suitable detection procedure, for example PCR. This method offers potential as an effective monitoring tool for emerging viruses and other pathogens by facilitating their rapid and sensitive detection. Therefore, the method described in the present study contributed to controlling current and future global infectious threats in the areas of food, medical and environmental science.

The efficiency of introducing amino groups onto the bead surface using the ammonia gas plasma technique may be increased by carefully adjusting the treatment conditions. Although the mechanism underlying the attachment of amino groups via plasma treatment remains to be fully elucidated, its clarification may further enhance the efficient introduction of these functional units. For example, beads with a higher number of amino groups are likely to adsorb a higher number of antibody molecules, resulting in the enhanced capture and concentration of viruses. In addition, the replacement of iron, which was used in the present study, with alternative metals possessing stronger magnetic properties for the fabrication of the beads may enhance their overall capture efficiency. Furthermore, a combination of various antibodies and beads may enable the multiplex detection of pathogens.

Following the initial emergence of DENV, its spread is enhanced in a time-dependent manner. Although vaccination strategies for DENV have not been established, it is likely that infectious viruses are indispensable for future vaccine production. Therefore, the viral isolation step is essential for vaccine production. Early and efficient vaccine production is particularly important during outbreaks of DENV-associated infections. Although the early detection and isolation of DENV from mosquitoes is crucial for preventing the potential spread of disease, progress towards the development of methods for DENV detection and isolation has been limited. Ultracentrifugation and PEG precipitation are conventionally used to concentrate viruses (8,9). However, these methods partially inactivate the viruses during the concentration procedure and are unsuitable for the routine monitoring of samples (8,9). The possibility of using magnetic beads coated with bioadhesive molecules to concentrate viruses has been suggested previously (25-42). An example includes anionic magnetic beads coated with an anionic polymer, poly (methyl vinyl ether-maleic anhydride), termed poly (MVE-MA), which can be used to concentrate DENV from infected mosquito cells derived from patients with dengue fever (37). Notably, the magnetic capture of other viruses using poly (MVE-MA) has been reported previously, including human immunodeficiency virus (35), borna disease virus (38), respiratory syncytial virus (39), influenza virus (40,41) and adenovirus (42). The most important aspects of this method are its simplicity and rapidity (<30 min). Furthermore, the applicability of antibody-integrated magnetic beads to the broad serotypes...
of DENV1-4 is another promising feature of this approach. Therefore, the method for DENV capture from solution, in combination with sensitive detection methods, may contribute to preventing the spread of different subtypes of DENV. In addition, the efficient capture of infectious DENV may assist in the development of vaccines against dengue fever. The magnetic bead-concentration method may facilitate the isolation and sensitive detection of DENVs, and may contribute to efficient surveillance and future vaccine production. The general utility of the method described in the present study may be further enhanced if it is found to be applicable to other emerging viruses and bacterial pathogens.

In conclusion, the present study demonstrated that antibody-integrated magnetic beads were useful for the capture of DENVs. The capture of DENV1-4 using the antibody-integrated magnetic beads was confirmed by the results of the RT-PCR analysis, showing that the BD fraction contained DENV genomic RNA. Therefore, this method may be used in combination with conventional PCR for the detection of DENV, and may increase the sensitivity of viral detection for the diagnosis of DENV.

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