Canine parvovirus (CPV) leads to an acute disease, characterized by hemorrhagic gastroenteritis, vomiting and myocarditis in dogs. The disease can affect dogs of any age but is fatal in pups. CPV has undergone genetic variations that have led to emergence of various CPV-2 antigenic variants such as 2a, 2b and 2c with replacement of the original CPV-2 circulating in the dog population. CPV genome is made up of 5.2 Kb nucleotides. Viral protein VP2 plays a very important role in determining antigenicity and host range specificity of CPV. The antigenicity as well as host range of CPV is determined by virus specific VP2 protein. That’s why the mutations that affect the VP2 gene are the main source of different antigenic variants. It spreads rapidly in the wild population of canines as well as domestic animals, infected feces serve as a main source of infection because the virus is shed in large quantity in the feces particularly 4 - 7 days post infection. The present review is focused on various
biotechnological approaches used for diagnosis of CPV along with some conventional techniques including gold standard virus isolation in animal cell culture, hemagglutination test, electron microscopy, enzyme linked immunosorben assay (ELISA). The biotechnological approaches such as polymerase chain reaction (PCR), Real-time-PCR, Loop-mediated isothermal amplification (LAMP), Bead based multiplexing, Microarray chip and DNA probe etc. have also assured their application. These approaches provide rapid, sensitive, optimal detection and effective control of CPV infection.

1 Introduction

Canine Parvoviruses (CPVs) are small, non-enveloped viruses belonging to the Genus Parvovirus in the family Paroviridae. Its genome is comprised of linear, negative-sense, single standard DNA of about 5.2 kb size, and encodes two structural (VP1 and VP2) and two non-structural (NS1 and NS2) proteins (Mochizuki et al., 1993). The first evidence of CPV infection in dogs dates back to 1970s and was identified as canine parvovirus type 2 (CPV 2) (Burtonboy et al., 1979). This CPV-2 isolate was likely to be a variant of Feline panleukopenia virus (FPV) because of detection of active circulation of intermediate viruses between FPV and CPV-2 in wild carnivores (Truyen, 2006). These two differed in at least six amino acids which are mostly located on VP2 protein (Truyen et al., 1995). Since then, CPV-2 has been identified globally and now, it is endemic in most populations of wild canines (Driciru et al., 2006, Ramsauer et al., 2007). Two new antigenic types of CPV-2. i.e. CPV-2a and CPV-2b differed at two amino acid positions, N426D and I555V (Truyen, 2006) and have became wide spread (Hoelzer et al., 2008).

The Asp-426 Glu substitution in capsid protein of CPV-2 generate a new variant known as CPV-2c which may infects several canine breeds (Buonavoglia et al., 2001, Decaro et al., 2006; Castro et al., 2007; Gombac et al., 2008). During acute phase of infection dogs may excrete virion particles up to 109/gram of faeces (Carmichael & Binn, 1981). Moreover, CPV-2 virion particles are very stable in environment which facilitates its transmission through faecal-oral route.

Canine parvovirus (CPV) infection is a highly infectious viral disease of dogs of great concern for pet owners, veterinarians and scientists due to its high morbidity and mortality rates associated. Parvovirus infects dogs of all age groups, but puppies are more affected than adults. The initial clinical signs of CPV infection are nonspecific and include anorexia, depression, lethargy, and fever. Within 24 to 48 hours, most affected dog starts vomiting and hemorrhagic small-bowel diarrhea results severe dehydration. With severe dehydration, protein loss, concomitant infection, and inability to produce a rapid immune response, further weakening the dog. In last, all of these factors can lead to shock and death (Bargujar et al., 2011). The current knowledge of epidemiology, pathogenesis, clinical findings and diagnosis of canine parvoviral enteritis was briefly highlighted (Geetha, 2015; Shim et al., 2015). CPV in faecal samples has been detected by several methods based on virus isolation in cell culture, haemagglutination (HA), electron microscopy, enzyme linked immunosorben assay (ELISA) and DNA hybridization.

Several molecular diagnostics assays such as polymerase chain reaction (PCR), multiplex PCR, Reverse Transcriptase PCR (RT-PCR), nucleic acid sequencing, Real Time-PCR, DNA probe etc have provided the unparalleled identification and discrimination ability for several viral pathogens (Minakshi et al., 2014; Kaur et al., 2015). Such diagnostic techniques should be transferred to end users for proper applications.

2 Conventional Methods

2.1 Viral Isolation

Various cell cultures viz. Crandell Feline Kidney cell line (CRFK), Madin Darby canine kidney cell line (MDCK), Walter Reed Canine Cell line (WRCC) have been used for the isolation of CPV from the clinical samples for diagnosis of canine parvovirus. CPV is primarily isolated in laboratory in canine lung and kidney cell line. It produces characteristics cytopathic effects (CPE) such as presence of intranuclear inclusion bodies in host cell, detachment and rounding of cells (Figure 1A, B). However, cell culture is not used as a routine diagnostic test because it is a time consuming process, require permissive cell lines, low sensitivity and requires skilled personnel. Moreover, the sensitivity of different cell lines for CPV multiplication may also vary. CPV-2 can be isolated from cell culture only after few days of inoculation (Desario et al., 2005).

2.2 Electron Microscopy

Electron microscope can also be used for morphological identification of CPV2. Under electron microscope CPV may be seen as either single virion particle or in group of few viruses (Amo et al., 1999). On 3rd to 9th day of infection viruses are excreted in large quantity in faeces, thus, this period is best for electron microscopic study of CPV-2 from fecal samples. However, electron microscopy needs large quantity of virus to confirm a sample as positive, because electron microscopy is less sensitive in comparison to other molecular tests (Esfandiari & Klingeborn, 2000). Electron microscopy was successfully used for diagnosis of canine parvoviral enteritis in fecal sample (Klingeborn & Moreno-López, 1980).
2.3 Haemagglutination (HA) Assay

Haemagglutination is specific, rapid and inexpensive test for CPV diagnosis. Haemagglutination is one of the important properties of Canine parvovirus. CPV has binding ability for sialic acid receptors on cell surface and agglutinates the RBCs of several animal species such as rhesus monkey, African green monkey, and Porcine etc (Burtonboy et al., 1979; Carmichael et al., 1980; Parrish et al., 1988). Seroprevalence studies among CPV2a infected dogs was reported using haemagglutination inhibition assay in North Korea (Klingeborn & Moreno-López, 1980; Deepa & Saseendranath, 2012). Two diagnostic assays was compared for their sensitivity and specificity and found that diagnostic accuracy of the ELISA was significantly greater than the IFA (Luren et al., 2012). However, the serology based diagnostic assays must be used with caution because in most of the cases dogs may show positive serological tests for CPV-2. This may happen due to administration of CPV-2 vaccine in large dog population and also the endemic nature of virus in several areas which may lead to inapparent infection and generation of antibody titer. Monoclonal antibodies based antigenic typing of canine parvovirus (CPV2a and CPV2b) was reported (Shankar et al., 2014).

2.4 Latex agglutination test

In latex agglutination tests the latex micro beads are coated with microbes specific either antigen or antibody, which can be used for detection of either microbial specific antibody or antigen in agglutination reaction. Positive result is detected by visualization of agglutination reaction which is characterized by clumping of micro beads with microbial antigen or antibody. Bodeus (1988) detected the CPV specific antibodies from field sample through latex agglutination test. Moreover, a new technology called SAT-SIT technology can be used for rapid detection of several other emerging hemagglutinating viruses from animals and humans (Marulappa & Kapil, 2009).

2.5 Counter immuno electrophoresis

A laboratory technique used to evaluate the binding of an antibody to its antigen. Counter immuno electrophoresis uses electric field in diffusion medium which is made up of polyacrylamide gel or agarose. The electric field facilitates the rapid migration of antibody and antigen towards each other so that line of precipitation will form at earlier than simple diffusion reaction. The line of precipitation indicates the binding of antibody with antigen hence positive result. Mixed infections for coronavirus antigen with canine parvovirus was detected by counter immuno electrophoresis in fecal samples (Ganesan et al., 1990). The prevalence of canine parvovirus infection was reported in clinically suspected dogs AGID and CIEP (Deepa & Saseendranath, 2012).

2.6 Fluorescent antibody test

In fluorescent antibody test an antibody tagged with fluorescent dye is used for detection of specific antigen. Based on tagging of fluorescent dye either on primary antibody or secondary antibody, the test may be either direct or indirect fluorescent tests. In direct fluorescent antibody test, the antibody binds directly with specific antigen and gives specific fluorescence signals for antigen detection. Fresh frozen tissues and formalin fixed are used to detect CPV using immunofluorescence (IFA) and immunoperoxidase (PAP). PAP gives more permanent, high resolution and clear intracellular localization of antigen than IFA (MaCartney et al., 1986). A semiquantitative ELISA and an immunofluorescence assay (IFA) were conducted for sensitivity and specificity of canine parvovirus (Gray et al., 2012). An indirect fluoroimmunoassay using magnetic protein micro bead was validated for identification of antibodies against canine viruses such as CPV, CDV and rabies (Wang et al., 2011).

Figure 1. A) Micro-photograph of CRFK Cell line after 48 hours of growth; B) Micro-photograph of CRFK Cell line showing CPE after 48 hours of infection.
Table 1 PCR assays developed for detection of canine parvovirus.

| S. No | Technique                                                                 | Gene/ target                                  | Reference          |
|-------|---------------------------------------------------------------------------|-----------------------------------------------|--------------------|
| 1.    | VP2 gene amplified by PCR and cloned in pTarget mammalian expression vector. VP2 gene was selected on the basis of restriction enzyme analysis and further confirmed by sequencing. The present work has shown that the recombinant plasmid could be used as DNA vaccine against canine parvovirus infection. | VP2 gene                                      | Gupta et al., 2005 |
| 2.    | Epidemiological study of canine parvovirus infection and analyzed by PCR assay. | sequences specific for CPV variant strains     | Behera et al., 2015 |
| 3.    | CPV-2a strains detected by PCR-RFLP                                       | VP2 gene                                      | Demeter et al., 2010 |
| 4.    | PCR was carried for VP2 capsid gene to detect all types of CPV, (CPV-2a/2b/2c) including the new CPV-2c | VP2 gene                                      | Silva et al., 2013 |
| 5.    | Genomic typing of canine parvovirus using PCR                              | sequences specific for CPV variant strains     | Costa et al., 2005 |
| 6.    | CPV and its variants typing using PCR                                       | sequences specific for CPV variant strains     | Shankar et al., 2014 |
| 7.    | Characterization of canine parvovirus by PCR                               | sequences specific for CPV variant strains     | Pereira et al., 2000 |
| 9.    | Analysis of VP2 gene sequences of canine parvovirus isolates               | VP2 gene                                      | Chinchkar et al., 2006 |
| 10.   | molecular characterization and phylogenetic analysis of canine parvovirus by PCR | sequences specific for CPV variant strains     | Mohan Raj et al., 2010 |
| 11.   | Typing of canine parvovirus using mini-sequencing based SNP analysis       | sequences specific for CPV variant strains     | Naidu et al., 2012 |
| 12.   | Detection of canine parvovirus by PCR assay                                | VP1 and VP2 gene                              | Singh et al., 2013 |

2.7 Enzyme Linked Immunosorbent Assay

The IgM antibodies indicate the recent infection of pathogen. These antibodies were derived in a number of laboratories; all appear to bind to the amino-terminal region of the major core protein. The sensitivity of ELISA tests is found to be much higher that other serological assays such as immunodiffusion test, HA or HI test (Banja et al., 2002). The sensitivity and specificity of sandwich ELISA for detection of CPV in dog fecal sample was found much higher than HA test (Rimmelzwaan et al., 1991; Drane et al., 1994). A point-of-care ELISA test kit yielded accurate results and highly sensitive and specific for detection of both CPV as well as CDV infection under field conditions. The Point-of-care ELISA system was used for identification of antibodies against CPV and CDV. This assay can be useful in animal vaccination programme and their care and management for outbreak of such disease (Litster et al., 2012). CPV antigens can also be identified in fecal samples by Sandwich ELISA (Deka et al., 2015).

3 Nucleic Acid Based Methods

Various nucleic acid based detection technique has been developed for the confirmation of CPV in the clinical samples. These techniques are fast, sensitive and specific and are discussed below:

3.1 Polymerase Chain Reaction (PCR)

PCR is a modern diagnostic assay which utilizes the specific amplification of desirable DNA sequence using template specific primer and DNA polymerase enzyme. It can also be used for diagnosis of those pathogens which may not be grown in laboratory condition. PCR assay has been used for diagnosis of several animal and human viruses. It can also detect viruses at early stage of infection before eliciting immune response and onset of clinical symptoms. Thus PCR may help in formulating policies for control and prevention of disease at early (Sharma et al., 2012). It can detect CPV from a samples having very minute quantity of virus. This assay is also much rapid and specific that gel filtration test. The samples having fecal inhibitory substances can be passing through spin column to remove inhibitory substances (Uwatoko et al., 1995). Molecular typing of CPV was done by using PCR based assays and CPV-2a and CPV-2b types were detected (Gauri et al., 2012). The PCR is a rapid, sensitive and specific method for detecting canine parvovirus (Savi et al., 2010, Figure. 2A, B). There are different researcher were used the PCR techniques for the diagnosis and detection of canine parvovirus as Table 1. Now a says several modifications of PCR such as multiplex PCR, Real-time PCR, nested PCR etc are used for molecular detection of viruses. Moreover, PCR amplicons can be used for nucleic acid sequencing and phylogenetic study for confirmatory diagnosis and tracking evolutionary history of virus.
3.2 Multiplex PCR

Multiplex PCR utilizes the power of PCR using several primer sets of different amplicons size for different pathogens in a single reaction. Multiplex PCR enables the presence of nucleic acids from several pathogens to be checked for in one test, but care must be taken to avoid interference between primer pairs or templates. It is a time as well as cost effective methods because it can detect several pathogens simultaneously. Multiplex PCR assay can also be used for simultaneously identification of canine Leptospira sp and CPV (Ramadass & Latha, 2005). The CPV-2a and CPV-2b strains were also differentiated using multiplex PCR assay (Parthiban et al., 2010).

3.3 Real-Time PCR

The real-time PCR is used for quantification of PCR product in reaction which can be used for estimation of viral load in sample. TaqMan assay based Real time PCR (RT-PCR) has been used for the detection of CPV-2 DNA in sample and an attractive tool for revealing single nucleotide polymorphisms in the capsid protein gene between CPV types 2a and 2b and CPV types 2b and 2c (Decaro et al., 2006). The advantage of the real time PCR is that there is no need to analyse the PCR product by agarose gel electrophoresis. Everything will be graphically shown on the monitor of the computer. Another advantage is that amount of the DNA present in the sample can be quantitated. Recently, SYBR Green based real time PCR has been developed for quantitation of CPV-2 variants in faecal samples of dogs (Kumar et al., 2010). Canine parvovirus infection was detected in feces of free-ranging wolves using real time PCR and the assay was 100% sensitive and specific with a minimum detection threshold level (David et al., 2012). RT-PCR method was used for the amplification of rotavirus RNA, BTV as well as for CPV viruses using Taqman probe and SYBR Green chemistry (Decaro et al., 2005; Anamul et al., 2015; Feng et al., 2015). The SYBR Green-based real-time PCR assay was used for the amplification of CPV 2, FPV and PPV DNA, with a reproducible limit of detection of as few as 10 copies/μL of target DNA per reaction and this study have been used successfully in veterinary diagnostic laboratory and have been helpful tools for the diagnosis and quantification of parvovirus infection in canines, felines and swine (Lin et al., 2014).

3.4 Multiplex Real-Time PCR

The term multiplex real-time PCR is used to describe the use of two to four fluorogenic oligoprobes for the discrimination of multiple amplicon. To date, there have been only a few truly multiplexed realtime PCR assays described in the literature. The use of non-fluorescent quenchers and the continuous development of better light sources in the machines are now in use and first applications for virus detection are becoming available. The vp2 gene based Multiplex Real-Time PCR was validated for simultaneously identification of CPV, FPV and PPV. Multiplex real time PCR have been used to detect and quantify CPV (Decaro et al., 2007; Wei et al., 2009; Zhao et al., 2013). Further, Kaur et al., 2016 reported that multiplex real time PCR assay could be used for rapid detection of CPV as well as typing of its three antigenic types.

3.5 PCR-Restriction Fragment Length Polymorphism (PCR-RFLP)

RFLP uses specific restriction enzymes for study of restriction pattern of viral nucleic acid. However, this is a time consuming technique. However, through PCR small quantity of viral nucleic acid can be amplified and used for RFLP analysis. The RFLP technique was successfully used for differentiation of CPV-2 antigenic variants (Savi et al., 2009; Zhang et al., 2010).
RFLP technique was also engaged in differentiation of CPV-2b and CPV-2c strains (Gauri et al., 2012). The partial VP2 gene specific PCR assay was standardized with corresponding consensus primers to amplify the desired length (747bp) of product. The PCR assay was carried out using the published partial VP2 gene specific primers (Sakulwira et al., 2001). Amplified 747 bp product was used for in silico restriction enzyme profiling. Reference sequences for vaccine and field strains were retrieved from the NCBI and loaded into the Insilco restriction enzyme profiling software (SNAP GENE SOFTWARE) & the resultant profiles were observed and images were obtained (Figure 3A). In wet-lab restriction enzyme profiling restriction enzyme RsaI (New England Biolabs) was used for template digestion. The different Restriction enzyme gave different RE digested products from both vaccines as well as field strain. The resultant digested products were resolved in 4% agarose gel electrophoresis (Figure 3 B).

3.6 Peptide nucleic acid-based (PNA) array

Peptide nucleic acid (PNA), are considered as a stable nucleic acid analogue. It contains pseudo-peptide skeleton in place of sugar phosphate backbone which is chemically and biologically highly stable. PNAs hybridize to cRNAs or cDNAs more efficiently than DNA. It possibly happens due to electrically neutral nature of PNA backbone. Peptide nucleic acid-based (PNA) array was used to discriminate between the four CPV-2 antigenic types (CPV-2, -2a, -2b, and -2c) during ante-mortem diagnosis of dogs, using newly developed PNA-DNA hybridization assay. The PNA array has high sensitivity and specificity compared with a real time PCR using the TaqMan assay, a gold standard method (An et al., 2012).

3.7 Nucleic acid hybridization assay

The VP1 and VP2 protein specific digoxigenin labeled probe have been developed for detection of CPV. This probe may also be used for in situ hybridization and detection of CPV from immobilized tissue samples in formalin and paraffins (Nho et al., 1997). Further, Decaro et al. (2005) develop two minor groove binders (MGB) fluorophores (FAM and VIC) labeled probe for rapid quantification of CPV-2 variants in dog fecal samples. The MGB probe was able to detect SNPs in CPV 2a/2b and 2b/2c. Both the MGB probe assays were found to be highly specific, sensitive and reproducible as compared to other methods used to detect the virus.

3.8 Loop-mediated isothermal amplification (LAMP) assay

LAMP assay is considered as alternative to conventional nested PCR. LAMP assay can be used for detection of DNA of several viruses of animal and human origin. In comparison to nested-PCR, LAMP assay are proved to be more rapid, sensitive and fairly reproducible method. It did not amplify other canine pathogens (Parthiban et al., 2012). Detection of canine parvovirus in fecal samples was reported using loop-mediated isothermal amplification (Cho et al., 2006). A detection system based on the application of LAMP in conjunction with ELISA and LFD for convenient visual detection of CPV with high sensitivity and specificity was developed (Sun et al., 2014). Mukhopadhyay et al. (2012) standardized a highly sensitive and specific LAMP assay for detection of CPV DNA from fecal samples. The assay showed result within one hour. Recently, VP2 gene based LAMP PCR assay has been developed (Figure 4). The assay is 30 times more sensitive than conventional PCR (unpublished data).
3.9 Whole genome amplification and sequencing

Complete coding sequence of canine parvovirus genome could be amplified by sequence specific primer. Pick-primer software is available in the NCBI BLAST to design these primers. To attain the maximal coding sequences overlapping of primers with their forward and reverse sequence is preferred. Optimization of PCR reaction mixture and thermocycling conditions is very important for amplification. The annealing temperature may vary with each primer and it should be standardized for the amplification. The products resolved in 1% agarose gel to observe the amplification (Figure 5).

3.10 Phylogenetic analysis

The nucleotide sequences of pathogens are used for phylogenetic study which shows its evolutionary relationship and closeness with other strains of same or different virus species. The phylogenetic studies of CPV vaccine strains in India have been done (Nandi et al., 2010). Phylogenetic studies also revealed the fact that Indian CPV variants are closely related among themselves. The CPV variants also showed little divergence from their ancestor MEVs (Singh et al., 2014).

The sequences were aligned against the other published CPV VP1/VP2 gene sequences using software from DNASTAR. The amino acid sequence, phylogenetic maps and percentage homology were deduced and analyzed from the sequences using the same software. The phylogenetic tree revealed that CPV2 and both CPV vaccine strains were in separate monophyletic group. The VP gene sequences of the Haryana isolate and gene sequences of various global isolates were used for phylogenetic analysis. The phylogenetic tree developed on the basis of VP gene indicated that the Hisar isolate is clustering with the Chinese isolate (Acc. No. JQ686671.1) as a separate group than rest of the Chinese isolates, Russian and USA isolates which indicates that the Hisar field isolate and the Chinese isolate originated from a common ancestor CPV (Figure 6).
3.11 Biosensor

Biosensor is an analytical device which converts biological responses to electrical signals. It is used for rapid diagnostic method which helps to detect disease in low sample with high selectivity and specificity in seconds. Disadvantage is that heat sterilization is not possible. A biosensor is also developed to detect CPV infection using Quartz Crystal Microbalance (QCM) biosensor and prolinker B to rapidly diagnose CPV infection. ProLinker™ B enables antibodies to be attached to a gold-coated quartz surface in a regular pattern and in the correct orientation for antigen binding. QCM biosensor is 95.4 % sensitive and 98% specific compared to PCR. It is rapid and accurate clinical diagnostic tool for CPV infection (Kim et al, 2015).

4 Prevention and control

The prevention and control of CPV infection depends primarily on an effective immunization program; but disinfection, animal movement control and husbandry practices also must be considered especially in shelters. In most of the cases of CPV treatment supportive therapy is used which is based on suppression of symptoms and prevention of further complications. Since disease is very acute the supportive intravenous fluid therapy should be started as soon as possible. The dog may recover within 2-3 days. However, the treatment may not always be successful. Care should be taken that infected dogs should not be allow to come in contact with other healthy dog, because CPV may infect other healthy dog easily.

The hygiene in kennel should be appropriate for disease prevention, because CPV can be live on some surfaces years' together. The bleaching solution in water in a ratio of 1:30 can be used to kill the CPV. Owner should not allow the dog to go outside and mix with other stray or infected dogs. Proper care should be taken in waste matter disposal especially faeces of infected dogs.

Conflict of interest

Authors would hereby like to declare that there is no conflict of interests that could possibly arise.

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