Molecular methods for rapid and specific detection of pathogens in seafood

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Seafood export are a major source of foreign exchange for many developing countries in Asia. Though generally seafood is considered a safe food, there are a few instances of outbreaks of food poisoning linked to seafood. Since seafood is often going to markets that are highly demanding in quality and safety, the seafood exporting countries need to take extreme care to ensure continued market access. One of the major concerns in seafood is the presence of pathogenic microorganisms. Because of this concern, often seafood processors have to ensure the absence of pathogenic microorganisms. Therefore quality assurance programmes require testing of raw material, product, water, ice and equipment for the presence of these pathogens.

Conventional microbiological methods for the detection of these pathogens is very time consuming and laborious. Figure 1 illustrates the general scheme for isolation and identification of pathogens. It takes between 4-5 days to detect pathogens by this method. Further problems arise due to situations such as the following:

Among V. cholerae bacteria, only serotypes O1 and O139 cause cholera. Among environmental isolates of serotype O1 non-toxigenic strains also exist.

Among V. parahaemolyticus only less than 2% of environmental strains are pathogenic. These are characterized by their ability to produce a thermostable direct hemolysin (TDH) or TDH- related hemolysin (TRH). While the former can be detected using a blood agar medium, Wagatsuma agar, there are quite often false positive reactions in this test. Preparation of this medium requires fresh (less than 24h) human or rabbit blood, which is difficult to obtain in most laboratories. There is no phenotypic test for TRH production.

DNA probe hybridization is based on the principle that (a) DNA is a double stranded molecule (b) the two strands of DNA can be separated by heating or chemical treatment (c) the two separated strands can reassociate (d) DNA strands from different sources can hybridise, provided there is complementarity of bases (A-T, G-C) between them.

Based on this principle, it is possible to make probes specific different microorganisms. Probes are short stretches of nucleotides that have sequences complementary to the target sequences. To detect probe hybridization, probes are labeled either with a radioactive molecule (p32), enzymes, ligands (eg biotin) or antigenic substrates (eg. digoxygenin). Figure 2 illustrates a typical probe.

Genes that are chosen as targets are those specific for each bacteria. In the case of pathogenic organisms, these are genes that encode virulence factors i.e. factors that make the organism pathogenic. Some examples of these genes are given in Table 1. Pathogenic V. cholerae produce a toxin, cholera toxin encoded by ctx gene. The virulence factors in V. parahaemolyticus are encoded by tdh and trh genes. The virulence factors in L. monocytogenes are encoded by iap and hly genes. In Salmonella, there are virulence-associated genes such as inv. Enterohemorrhagic E.coli have virulence genes such as stx, eae. Using such specific probes, it is possible to specifically detect the pathogenic strains of bacteria. Figure 3 shows a general protocol for detection of pathogenic bacteria using colony hybridization.
The food sample is homogenized (1:1) in a buffer and plated on a non-selective agar. The plates are incubated at 35°C for 18 h. The colonies appearing on the plate are transferred to a suitable filter. The bacterial cells are lysed by immersion in a lysing solution and the DNA released are denatured and fixed to the filter.

The filter is now incubated in a prehybridising solution and then the probe is added. Hybridisation is performed at a temperature appropriate for the probe. After hybridization, the filters are washed at a specified temperature. Probe hybridizing to DNA on the filter is detected depending on the type of label. If radioactive probe is used, detection is by autoradiography, where in the filter is incubated with a X ray film, one can see spots corresponding to colonies to which probe has hybridized.

However, presently a number of non radioactive probe labels are available. Most convenient are enzyme labels which can be detected using the appropriate chromogenic substrate. Figure 4 shows colony hybridization for detection of *V. parahaemolyticus* in seafood using an enzyme labeled probe.

To detect pathogens that may be present in extremely small numbers, the food samples may be enriched before plating. The filters after hybridization can be preserved as a record of analysis. Probe hybridization analysis requires no sophisticated equipment. A hybridization incubator is all that is needed. Therefore this technique can be most conveniently adopted in seafood quality control laboratories. In some situations, DNA probe-based methods are essential to detect pathogenic strains of organisms eg. *Vibrio parahaemolyticus*. This organism is commonly found in coastal and estuarine areas all over the world. 98% of environmental strains are not pathogenic. Hence mere detection of this organism by conventional microbiology is not sufficient to determine the hazard. DNA probe hybridisation methods are getting wide acceptability in quality control laboratories and the US Food and Drug Authority Bacteriological Analytical Manual describes this method indicating acceptance by the regulatory agencies. In the case of bacteria such as *Salmonella* and *Listeria monocytogenes*, DNA probe hybridisation methods have undergone multilaboratory evaluations conducted through AOAC and are accepted as official methods.

**Polymerase chain reaction and its applications in assessment of seafood safety**

PCR is a nucleic acid amplification technique wherein a specific portion of nucleic acid from a target organism is amplified in vitro. This specific amplification is achieved using oligonucleotide primers that are specific for the region flanking portion to be amplified. The amplification requires the enzyme DNA polymerase, and the building blocks of DNA, the deoxyribonucleotides (dATP, dTTP, dGTP, dCTP). The reaction is performed in several cycles, each cycle consisting of three steps (a) DNA denaturation: this is the step in which the target DNA strands are separated by heating to about 95°C. (b) Primer annealing: this is the step in which the primer binds to the target region specifically. This step is carried out at 55-65°C (c) Primer extension: this is the step in which the new DNA strand is synthesized by the DNA polymerase on the template strand. Normally about 30 cycles of reaction are performed. Since each cycle involves denaturation of DNA at 95°C, the DNA polymerase used in the reaction should be thermostable. The discovery of thermostable DNA polymerase from the thermophilic bacterium *Thermus aquaticus* led to rapid application of PCR in diagnostics.

By designing oligonucleotide primers that are specific for an organism, it is possible to design PCR to amplify specifically DNA from any desired organism. In the case of RNA viruses, it is possible to first copy the RNA into DNA using the enzyme reverse transcriptase.

**Table 1: Examples of pathogens and target genes**

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Target genes for probe hybridization</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Vibrio cholerae</em></td>
<td>ctx</td>
</tr>
<tr>
<td><em>V. parahaemolyticus</em></td>
<td>tdh, trh</td>
</tr>
<tr>
<td><em>V. vulnificus</em></td>
<td>Vvh</td>
</tr>
<tr>
<td><em>Salmonella</em></td>
<td>inv, hms</td>
</tr>
<tr>
<td>Enterohemorrhagic E.coli</td>
<td>stx, eae</td>
</tr>
<tr>
<td>Enterotoxigenic E.coli</td>
<td>St, Ct</td>
</tr>
<tr>
<td><em>L. monocytogenes</em></td>
<td>iap, hly</td>
</tr>
</tbody>
</table>

**Figure 4. Detection of total *V. parahaemolyticus* using probe for thermolabile hemolysin and pathogenic V. parahaemolyticus using probe for thermostable direct hemolysin (B)**
transcriptase. PCR used to detect RNA targets is referred to as RT-PCR.

PCR for detection of pathogenic bacteria in seafoods

PCR technique for detection of most pathogenic bacteria associated with seafood have been described. In most cases the oligonucleotide primers have been designed to specifically amplify virulence associated genes. For example contamination of seafood with toxigenic V.cholerae can be detected using PCR amplifying the ctx gene encoding the production of cholera toxin.

“98% of environmental strains are not pathogenic...”

Contamination of seafood with pathogenic V. parahaemolyticus can be detected using PCR amplitifying the tdh and trh genes that encode virulence associated hemolysins. In the case of pathogenic Escherichia coli, the potential targets for amplification include stx gene encoding the production of shiga-like toxin, eae gene encoding intimin, heat-labile (LT) and heat stable toxins (ST) etc. In the case of Listeria monocytogenes, several target genes have been reported. These include the gene encoding the production of the invasion associated protein, iap, listeriolysin, hlyA, and the regulatory protein, prfA.

PCR is a DNA amplification technique and therefore, even if there are dead bacteria, they would show up in PCR. Therefore for determination of seafood safety, it would be important to ensure that only viable pathogens are detected. This can be achieved if PCR is performed after enriching the food sample in suitable broth.

“...using specific probes, it is possible to specifically detect the pathogenic strains of bacteria.”

PCR Based techniques for tracing the source of contamination

Traditional methods of identifying bacteria by biochemical tests can identify them up to species level, but cannot differentiate between strains. Techniques such as serotyping and phage typing also have little discriminatory power. PCR based techniques such as Random Amplification of polymorphic DNA (RAPD) can generate DNA fingerprints of organisms. RAPD patterns are helpful in studying similarity or differences in strains. For example, this technique has been used to differentiate strains of Listeria monocytogenes isolated from raw fish and from smoked fish and from processing environments. This type of study would help in understanding the source of the strains found in the product, eg cold smoked fish. In this technique single 10 mer oligonucleotide primer is used to perform amplification at low annealing temperature (eg 37°C). The primer is not targeted at any particular region of the genome and therefore, this reaction can be performed even in the case of organisms whose genome sequence is not available. Figure 5 shows an illustration of the application of this technique. In our laboratory, we have been using RAPD to study the source of contamination of pathogenic Vibrio spp in seafood.

Conclusion

The safety of products of aquaculture has been a concern and assessment of safety requires detection of pathogenic microorganisms. Conventional methods for detection are time consuming and cannot discriminate between pathogenic and non-pathogenic strains. Molecular techniques such as PCR and colony hybridization are useful for rapid detection of pathogens and specific detection of virulent strains. Since these are rapid, specific and sensitive, they have immense applications in seafood quality control laboratory.

References