Introduction

Foodborne illnesses are usually caused by the ingestion of food products contaminated with pathogens or toxins. Yersiniosis is the third-largest foodborne disease in the European Union instigated by Y. enterocolitica, which is a Gram-negative coccus belonging to the Enterobacteriaceae family, where it shares about 10–30% DNA homology with other family members. Biochemically and serologically, it has been categorized into six biovars, 70 serotypes and three species (Y. enterocolitica, Y. pestis, Y. pseudotuberculosis), which are responsible for foodborne infections in humans.

Yersiniosis is self-limiting gastroenteritis, with severe invasive sickness like terminal ileitis, mesenteric lymphadenitis, mimicking appendicitis, and sometimes septicemia. Furthermore, post-infections in immunocompromised patients lead to erythema nodosum, arthritis, and glomerulonephritis. The CDC estimates about 1 177 000 cases of Yersiniosis annually in the US caused by Y. enterocolitica infections, and 90% of these are foodborne. The diverse geographical distribution is the primary cause of the global epidemic of Yersiniosis, as shown in Fig. 1.

The transmission of Y. enterocolitica occurs during blood transfusions, through which it causes the highest cases of mortality. The primary sources of this bacterium are animal-originated foods, especially pork and pork products. The propagation of Y. enterocolitica at low temperatures has marked it as a life-threatening bacterium; hence, it has been linked with foodborne infections. Some studies have revealed that it can also survive and reproduce in vacuum packaging and refrigerated foods. Food-based Yersiniosis epidemics are linked with multiple food items, including dairy, meat, poultry, fruits, vegetables, stewed and fermented products and also seafoods, as given in Fig. 2. The infective dose of Y. enterocolitis is about 10^3 to 10^6 CFU mL^−1 for human infections, which has not been investigated in previous studies.

In 1976, the first outbreak of Yersiniosis was reported in New York due to the consumption of Y. enterocolitica-contaminated chocolate milk. Similar outbreaks of milk-borne Yersiniosis were also reported in other parts of the United States, Europe, Australia, Sweden and India. Subsequently, several outbreaks of Yersiniosis caused by eating contaminated pork products were reported in Hungary. Subsequently, several outbreaks of Yersiniosis caused by eating contaminated pork products were reported in Hungary, United States, Norway, China, and European countries. Frequent cases of Yersiniosis in European countries are rarely associated with beef, mutton, milk and dairy products; therefore, the distribution of Y. enterocolitica is highly diverse as it can contaminate both animal and
plant-based foods, thus triggering an alarming situation of foodborne illness for the food authorities. Yersiniosis has been reported several times in the past few years around the globe; however, the source of infection has remained unknown\(^2\) in most of these cases.

\textit{Y. enterocolitica} can survive and grow in contaminated foods, both at the industrial and commercial levels. The latest trend of the increased consumption of processed foods has further enhanced the risk of Yersiniosis. Moreover, the global emergence of Yersiniosis is also linked with the growth of international food trade, the revolution of animal husbandry and the development of the food processing industry.\(^2\) The Foodborne Diseases Active Surveillance Network (Food Net, USA) studied the emergence of Yersiniosis for about ten years (1996-2007) and concluded that the cases of Yersiniosis in the United States were miscalculated due to the lack of detection methods. On the other hand, in a few other countries, food samples are not considered for \textit{Y. enterocolitica}, which indicates that the diagnosis is inadequate.\(^2\)\(^5\)\(^3\) A timely and adequate detection of \textit{Y. enterocolitica} in food samples is highly recommended in order to prevent the distribution of this pathogen. Therefore, updated information about the morphology, classification, sources of infection, pathogenicity, and cultural and molecular detection methods is summarized in this review, which will serve as a guideline for researchers and highlight the key research areas for the development of rapid detection methods.

### Sources, classification, and biology

\textit{Y. enterocolitica} is a rod-shaped facultative anaerobic, non-spore-forming Gram-negative bacterium habitually found in nature, including food, water, and animals. However, pigs are considered as the main reservoirs.\(^3\) It is estimated that 35–70\% of swine herds and 45–100\% of individual pigs carry \textit{Y. enterocolitica}. According to pathogenicity and geographical distribution, it is categorized into five distinct groups: 1A, 1B, 2, 3, 4, 5. \textit{Y. enterocolitica} has about 60 serotypes. Some serogroups belong to each biotype as follows: 1A (O:5; O:6, 30; O:7, 8; O:18; O:46), 1B (O:8; O:4; O:13a, 13b; O:18; O:20; O:21), 2 (O:9; O:5, 27), 3 (O:1, 2, 3; O:5, 27), 4 (O:3) and 5 (O:2,3).\(^2\)\(^2\)\(^3\)\(^3\) The predominant serogroups that cause the most infections are O:3, O:8, O:9, and O:5.\(^5\)\(^3\)\(^4\) Some of the biochemical features are described in Table 1.

The biphasic lifestyle of \textit{Y. enterocolitica} allows it to grow within or outside the human body. It produces pinpoint colonies and shows slow growth on sheep blood agar, MacConkey agar, and Hektoen enteric-coated agar plates. The selective isolation was reported on \textit{Y. enterocolitica} cefsulodin-Igansanovibomycin (CIN) and virulent \textit{Y. enterocolitica} (VYE) agar plates. \textit{Y. enterocolitica} can grow at both low and high temperatures ranging from 0 to 44 °C, but 25–30 °C is considered as the optimum growth range. The growth temperature also affects the physiology of the bacterial cells. A study revealed that at lower temperatures, it becomes motile due to the growth of peritrichous flagella and more virulent due to the transcription of virulent factors like ystA, enterotoxin production gene \textit{ail}, and invasion of \textit{locus gene}, \textit{rfbC}, and \textit{O-antigen genes},\(^3\) while at higher temperatures, it becomes non-motile and lacks flagella. Besides, it can even grow at 1 °C in milk and raw meat in a 5%
sodium chloride solution and at pH up to 4.6 (ranging from 4 to 10). Thus, the growth ability in extreme conditions makes it a more alarming pathogen that can multiply in different conditions and therefore, it is considered as a severe threat to the consumers.

Virulence factors and pathogenic mechanism

Refrigerated contaminated food products, e.g., chocolate milk can only result in Yersiniosis, if they carry high doses \(10^7 - 10^9\) CFU mL\(^{-1}\) of \(Y.~enterocolitica\) cells. The bacterial cell starts infecting the small intestine at terminal ileum and proximal colon, which are considered as the primary sites for infection. It is presumed that bacteria mainly use virulent gene products to colonize inside the intestine and with the rise in temperature \(37~^\circ C\), they induce the expression of virulent gene \(pYV\) for the secretion of virulent factors. Later, these virulent factors (ail,
Table 2  *Y. enterocolitica* key virulence factors and their genetics

| Origin                  | Sizes      | Function                                                                 |
|-------------------------|------------|--------------------------------------------------------------------------|
| Chromosomes             |            |                                                                          |
| Invasin                 | 92 kDa     | Inv binds to β1-integrin and promotes adhesion and invasion             |
| Attachment invasin locus (Ail) | 17 kDa    | Attachment and invasion: serum resistance                                |
| Yst (enterotoxin)       | 3.5–6 kDa | Yst increases cGMP level and fluid secretion                              |
| Yersiniabactin (catechol-tye) | 482 kDa   | Siderophore, an iron-binding protein                                      |
| Virulence plasmid (pVY) | 70 kb      |                                                                          |
| Ysc (Yop secretion)     | 28 proteins| A type III secretion system (TSS)                                        |
| YopH                    | 51 kDa     | Dephosphorylated host proteins modulate signalling pathways and prevent phagocytosis |
| YopD                    | 33.3 kDa   | Responsible for translocation of YopE and other effector proteins (YopH, YopM, YopO, etc.) across the membrane |
| YopE                    | —          | Inactivates Rho family of GTPase, as a result, disrupts actin cytoskeleton and prevents phagocytosis |
| YopP                    | 33 kDa     | Macrophages apoptosis; alters the expression of cytokines                |
| YopT                    | —          | Interferes with actin cytoskeleton formation by inactivating Rho GTPase |
| YadA (adhesion protein) | 160.240 kDa| Adhesion to epithelial cells by interacting with 1-integrin: blocks complement-mediated killing; serum resistance |
| YopB                    | 41.8 kDa   | Inhibit cytokine release from macrophages                                |
| LerV                    | 37.2 kDa   | Low calcium response                                                     |

*Source: Foodborne Microbial Pathogens (*Yersinia enterocolitica* and *Yersinia pestis*).*37

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**Fig. 3**  Mechanism of the pathogenesis of *Y. enterocolitica*. The translocation of *Yersinia* is done via intestinal epithelial cells. Afterwards, they enter the basal layer through M-cells and start invasion through the interaction with the β1-integrin of the host cell. Later, they are transported to the lymph nodes and the liver by macrophages/dendritic cells, where they start delivering YOPs via type III system (TTSS) into the cytosol of the host cell. *Y. enterocolitica* safeguards itself from phagocytosis and macrophage apoptosis by retarding cytokine production.
Yop unnecessarily facilitate the binding of the bacterial cells to the mucous membrane and invade M-cells. Invasive proteins interact with β1-integrin receptors, which are abundantly located on M-cells. The virulent factor YadA supports the invasion by interacting with β1-integrin receptors, collagen, fibronectin, and laminin. Then, the trapped bacteria are freed from the M-cells in the basal layer of the lamina propria and after multiplication in lymphatic follicles, they cause necrosis and abscesses in Peyer’s patches.55

The key virulence factors are shown in Table 2. Besides, bacteria can reinvoke the epithelial cells by using the β1-integrin receptors positioned of their lateral basal surface. Bacteria spread from Peyer’s patches to mesenteric lymph nodes, causing typical lymphadenitis.5, 34 The bacterial cells also spread to the liver, spleen and lungs and propagate by protecting themselves from the phagocytosis of polymorphonuclear macrophages such as leukocytes and neutrophils. YOP is transmitted to the macrophages by TTSS, which blockades the process of phagocytosis and the oxidative burst of phagocytosis and induces apoptosis to interfere with cell signal transduction events, thus promoting the survival of the bacterium. Yersinia also inhibits inflammation by preventing macrophages and other immune cells from releasing pro-inflammatory cytokines (TNF-alpha, IFN-gamma, IL-8). An overview of the pathogenesis mechanism is given in Fig. 3. In general, the TTSS system halts phagocytosis and suppresses the immune system, thus safeguarding the survival of bacteria in lymphoid tissues.

Furthermore, they cause abscesses in Peyer’s patches by secreting enterotoxin Yst (to promote cell fluid secretion) to damage the epithelium; Yst-a, Yst-b and Yst-c activate membrane-bounded guanylate cyclase, resulting in the bigger deposition and activation of cyclic guanosine monophosphate (cGMP) in cells, followed by the stimulation of cGMP-dependent protein kinase. Enterotoxins terminate the biological reactions that inhibit sodium absorption and stimulation of chloride secretion. The resulting adverse effects can lead to septicaemia, pneumonia, meningitis, and endocarditis, which can be lethal to immunocompromised hosts or individuals with underlying diseases. Y. enterocolitica infections can also develop nosocomial infections, arthritis, diarrhoea and sequelae in some patients.15, 17, 48 The understanding of the pathogenic mechanisms of this bacterium will help us develop strategies for the prevention and control of Y. enterocolitica.

Isolation and detection

The second part will present the available isolation and detection strategies for this pathogen. Several Yersiniosis cases reported in the past two decades have emphasized the necessity to study this pathogen and develop rapid, accurate, and economical detection methods. To prove the relationship between pathogens and epidemics, it is necessary to isolate and identify pathogens from suspicious food products. This is a challenging task because this bacterium grows slowly and dominates in the sample.7 Some phenotypic or molecular characteristics that are used to identify pathogenic strains rely on the presence of a virulent plasmid (pYV), while other characteristics are lost during the prolonged processes of culturing, isolation and detection. The in situ detection of Y. enterocolitica requires the separation of bacterial cells from the raw sample and the subsequent detection by different cultural methods.59 These methods are usually applied for qualitative analysis, but certain limitations like long latency, low sensitivity and lack of differentiation between pathogenic and non-pathogenic strains reduce the applicability of these methods. Thus, new formulations and standardization of these methods are still required for Y. enterocolitica.

Cultural methods

For decades, cultural methods have been used as the primary standard methods for pathogen detection. They can be applied in the detection of pathogens from natural sources like water, food and clinical samples. However, the high costs, time-consumption, complexity, and low sensitivity of these methods hinder their applicability in the routine diagnosis of pathogens. Additionally, the concentration of Y. enterocolitica in the samples is always below the detectable limit, which requires additional pre-concentration steps to apply these culture methods for diagnosis.

In many studies, different selective media for pre-concentration have been introduced to solve this problem. Selective media have been widely used for pre-enrichment with significant advantages; for example, an irgasan-ticarcillin-potassium chloride (ITC) broth was used to increase the number of bacteria. Similarly, other formulated media like MacConkey agar, Hektoen enteric (HE) agar, xylose lysine deoxycholate (XLD), cefsulodin irgasan novobiocin (CIN) agar and virulent Y. enterocolitica (YVE) agar have been used for the isolation of Y. enterocolitica. CIN agar and esculin-modified CIN agar are used as differential media, on which this bacterium produces red “bull’s eye” colonies, as shown in Fig. 4, whereas other Yersinia spp. produce dark colonies.49 Phosphate buffer saline (PBS) and trypsin soybean soup (TSS) have been reported as the best media for the pre-enrichment of the samples of Y. enterocolitica.77, 78, 41

However, in the culturing methods, the prolonged incubation time leads to the growth of other psychrophilic bacteria, which limits their further application. This problem was overcome by employing an alkali treatment to enrich the culture in Y. enterocolitica and reduce the competitive background microflora since Y. enterocolitica can multiply in an alkaline environment as compared to other Gram-negative bacteria.43 In another study, selective media containing discriminatory antimicrobials were formulated to concentrate the bacterial cells at high temperatures like the Modified Rappaport Broth (MRB) applied for the rapid recovery of serotype O:3 at 25 °C for 2–4 days and for the quick recovery of bacterial cells, the selection of media depends on sample types. Like selenite medium was used for meat samples, irgasan-ticarcillin-chlorate (ITC) broth for isolation of serotype O:3, improved TSB for recovery of Y. enterocolitica from ready-to-eat food and pork samples. Luria–Bertani bile salt irgasan (LB-BSI) was also supplemented with
Cefsulodin and novobiocin for the recovery of *Y. enterocolitica* at 12 °C. The above media are used for the pre-concentration of *Y. enterocolitica* and other selective media were reported for its isolation, including MacConkey agar, Salmonella Shigella deoxycholate calcium (SSDC) agar and Cefsulodin-Irgasan-Novobiocin (CIN) agar. Although three different chromogenic media, namely, *Y. enterocolitica* chromogenic medium (YeCM), *Y. enterocolitica* agar-selective chromogenic medium (YECA) and CHROMAagar (CAY) have also been developed for use in diagnostic and clinical laboratories, it was observed that CIN Agar modified with L-arginine, ferric ammonium citrate, sodium thiosulfate and DL-phenylalanine had better identification ability for *Y. enterocolitica* over other bacteria in food samples. Importantly, the ISO method for food samples (ISO 10273:2003) also includes the use of the PBS broth, ITC broth for pre-concentration, and CIN for plating. Similarly, other developed methods, such as NMKL method No-117, NCFA, BAM-U.S. FDA, indicated the cold enrichment of *Y. enterocolitica* on PSB and MRB. These expensive and labour-intensive laboratory preparations persist as a challenge for the routine screening of *Y. enterocolitica* in the food industry. Therefore, further improvements are needed in the traditional culture methods for the isolation and identification of *Y. enterocolitica*.

**Immunological methods**

The recent advancements in pathogen detection have improved accuracy and time. In this context, a lot of immunoassays have been reported for the quick recognition of microorganisms without culturing. The simplest method developed for the detection of *Y. enterocolitica* serotype O:3/O:9 and serotypes O:3, O:6 or O:9 involves the use of latex agglutination assays, in...
which specified antibody-coated latex beads are agglutinated with antigens to form visible precipitates.29 The immunomagnetic separation (IMS) strategy reduces the detection time by 24 hours and eliminates the pre-enrichment step. In this method, magnetic beads covered with target-specific antibodies are used to sort out the pathogen from a cocktail of microorganisms, as shown in Fig. 4. In one study, Y. enterocolitica could be sorted out by the IMS method and confirmed by nested PCR using the specified primers for the Y. enterocolitica adhesion gene (yadA), achieving $10^4$ to $10^5$ CFU mL$^{-1}$ LOD (limit of detection) for clinical and environmental samples.30 This method was more specific and could differentiate pathogenic and non-pathogenic strains without any cross-reactivity. However, this method was applied to detect pathogenic strains lacking the pYV virulent gene. A new surface plasmon resonance (SPR)-based immunosensor constructed for the quick recognition of Y. enterocolitica with LOD of $10^5$–$10^7$ CFU mL$^{-1}$ was developed, where the antibody–antigen reaction was used for detection by immobilizing the antibodies on the SPR surface.31 A sandwich-type multiplex chemiluminescence immunoassay was established by using peroxidase-labelled monoclonal antibodies; the peroxidase activity of bound polyclonal antibodies was measured by an enhanced luminol-based cocktail using a low-light charge-based coupled imaging device. This method is considered to be rapid and accurate for detection as compared to the conventional culturing methods. It can detect up to $10^5$–$10^7$ CFU mL$^{-1}$ bacterial cells in samples with high recovery values of around 90–120%.32 Direct antigen detection provides rapid and specific identification results; however, the specificity of these methods is limited by the quality of the antibodies. These analyses usually require many antigens in the sample, which involves concentration steps before analysis, thus increasing the detection time. Also, these

Table 3 Recently developed detection methods for Y. enterocolitica

| Country     | Food                               | Method                                                                 | Detection time | Advantages/disadvantages                                      | Reference |
|-------------|------------------------------------|------------------------------------------------------------------------|----------------|--------------------------------------------------------------|-----------|
| South Korea | Kimchi                             | Single-walled carbon nanotube (SWCNT)-based biosensor                 | >a day         | Pre-enrichment                                              | 68        |
| Finland     | Milk and human serum               | GQD-based immunosensor Cultural & real-time PCR (ail genes)           | >a day         | Rapid, specific                                              | 69        |
|             | Raw milk, Iceberg lettuce, minced meat | SNP analysis (ail, inv, virF, yadA, ystA, ystB, myfA, irp2, fyuA) | 5–10 days     | Expensive instrumentation, trained staff                   | 48        |
| China       | Various foods                      | Duplex PCR combined with capillary electrophoresis laser PCR (ail, ystA, ystB, and inv) | >a day         | Enrichment not required                                      | 70        |
| Italy       | Pigs and piglets                   | Real-time PCR (ail gene)                                              | 2–3 days       | Expensive instrumentation, trained staff                   | 71        |
| Italy       | Milk                               | SNP analysis (ail, inv, virF, yadA, ystA, ystB, myfA, irp2, fyuA)      | 3–4 days       | Pre-enrichment is required                                  | 41        |
| Poland      | Pig tonsils                        | Real-time PCR (ail gene)                                              | 2–3 days       | Expensive instrumentation, trained staff                   | 72        |
| France      | Pigs                               | Pulsed-field gel electrophoresis (PFGE) and MLVA                     | 1–2 days       | Expensive instrumentation, trained staff                   | 73        |
| Italy       | Meat, dairy products               | Captured-ELISA (46F7, 54 B11, 54C 11, 62D10, 6 4C7, 6 4C10, 72E8, 72E10, 72G6) | 1–2 days       | Trained staff                                               | 8         |
| Brazil      | Human source                       | Duplex PCR tufA and rfbC                                              | Expensive instrumentation, trained staff               | 74        |
| Czech Republic | Fruits and vegetables               | Real-time quantitative PCR (qPCR) without pre-enrichment (ail gene) | 1 day          | Expensive instrumentation, trained staff                   | 1         |
| Poland      | Human samples                      | ELISA MyfA and PsaA                                                  | Not specified  | Expensive instrumentation, trained staff                   | 75        |
| Poland      | Pig tonsils                        | Cultural plus real-time PCR (chromosomal locus tag CH49-3099 gene)     | Not specified  | Expensive instrumentation, trained staff                   | 72        |
| Germany     | Pork meat                          | In situ hybridization (16S and 23S ribosomal RNA)                     | 1 day          | Expensive instrumentation, trained staff                   | 76        |
| Poland      | Pig tonsils                        | PCR and culture methods (inv- positive and ail-positive)             | Not specified  | Expensive instrumentation, trained staff                   | 77        |
| Denmark     | Blood samples                      | LPS-based ELISA (lgA, lgG, and lgM)                                  | Not specified  | Diagnostic only                                             |           |
| Canada      | Mix food items                     | Multiplexed real-time PCR (qPCR) and pyrosequencing                   | Not specified  | Expensive instrumentation, trained staff                   | 78        |
immunization methods may be affected by in vitro test conditions and may lead to false detection.

Molecular techniques

The molecular technology has also been considered as a promising tool for the quick detection of pathogens by limiting the problems of the culture technology. The culture technology is time-consuming and laborious and the identification results of pathogenic and non-pathogenic bacteria are also unreliable.\(^\text{29}\) Genomic targets are used in molecular techniques for the specific detection of pathogens in food samples. Standard methods include colony hybridization, PCR, isothermal amplification and microarray technologies. Generally, the polymerase chain reaction (PCR) is considered the most suitable technique for the detection of *Y. enterocolitica*. In this review, we have summarized the latest molecular and immunological detection methods (Table 3).

Molecular typing methods

Pathogens always cause severe illnesses, but their detection in food samples still remains a challenge. In this regard, nucleic acid-based detection analyses are very prominent. The revolution in the molecular technology has enabled scientists to develop a series of fast, economical, and accurate molecular techniques. Different examples include population hybridization, polymerase chain reaction, nested polymerase chain reaction, real-time polymerase chain reaction, microarray technology, and loop-mediated isothermal amplification (LAMP) technique.\(^\text{54}\) Commonly applied methods for the detection of *Y. enterocolitica* include the probing of target virulence-related DNA sequences, such as *ail* (attachment and invasion locus), *inv* (invasin), or *yst* (*Yersinia* stable toxin) genes.\(^\text{55}\) Besides the great advantage of the higher sensitivity of molecular methods over traditional cultural methods, there are still certain limitations, like the competing microflora, which tend to decrease the hybridization efficiency.

In the field of microbiological diagnostics, PCR-based assays have gained great popularity due to their speed, cost-effectiveness, and the comfort of automation. In most cases, the polymerase chain reaction (PCR) uses chromosomal virulence genes, such as *ail*, *inv*, and *yst*, as the targets or main targets of *Y. enterocolitica* detection. The limitations of pre-enrichment, pre-PCR sample preparation, selection of the right single gene or a mixture of genes, and analysis of PCR products result in the establishment of new types of PCR assays. The European Committee for Standardization’s (CEN) recommendations regarding the PCR-based detection protocols of foodborne pathogens have led to the development of real-time PCR-based methods. Real-time PCR meets the criteria of CEN due to its high detection probability, good accuracy, simplicity, little or reduced residue contamination and easy-to-access protocols. Recently, the real-time polymerase chain reaction (PCR) has been applied to detect *Y. enterocolitica* in food samples. These methods use virulence-related genes, especially *ail* genes. Though these PCR-based methods meet the criteria of CEN, some issues arise like the fact that common molecular markers (*ail*, *myf*, *yst*) on the pathogenic and non-pathogenic bivars of *Y. enterocolitica* make it hard to differentiate viable and non-viable cells, while the presence of inhibitory substances in food samples limits their application. One way to resolve these issues is by introducing pre-enrichment steps before PCR. The use of negative controls limits cross-contamination for post-PCR measures. Moreover, PCR can be multiplexed to allow multiple detections of targets to ensure high specificity. Several multiplex PCR methods have been developed to amplify more than two virulence genes for the specific detection of *Y. enterocolitica* in food samples.\(^\text{56}\) Weagant and co-workers developed multiplex PCR using rfbC (O-antigen encoding gene), *inv*, *ail*, and *virF* gene primers for the specific detection of *Y. enterocolitica* serotype O:3.\(^\text{57}\) Recently, a new PCR-based approach was developed for the identification and serotyping of *Y. enterocolitica*, where the first multiplex PCR identified species and subspecies levels using 16SrRNA, *inv*, *ail*, and *ystB*, followed by nested PCR for differentiating the serotypes associated with human infections using serotype-specific primers, rfbC/wbbU/wbaA/wzt.\(^\text{57}\) As most of these are based on “Taqman and SYBR green” methods, this gene-based serotyping scheme can be applied for the reliable identification of toxigenic *Y. enterocolitica*.\(^\text{58,59}\)

The high-resolution melting technology (HRMT) is one of the recent advancements in molecular biology for the detection of bacteria by single nucleotide polymorphisms (SNPs), hyper-variable repeats and mutations in PCR amplicons. The changes in nucleotides at specific sites can be monitored in closed-tube analysis after PCR and then, the fluorescence of the PCR products labelled with specific dyes can be detected to observe the transformation of unfused and melted DNA. Although, this method reduces the time and minimizes the risk of cross-contamination, it requires sequence variants with desired melting points and labelled primers.\(^\text{60}\)

Although, PCR techniques have many benefits over traditional culture methods, they still require highly trained labour, expensive instruments (electrophoresis and thermal cycler machines) and complicated sample pre-treatments, which make their application difficult for real samples. Lately, a new technique of isothermal amplification has been introduced, which simplifies the amplification of targeted nucleic acids at the laboratory level with inexpensive instrumentation, requiring only a water bath at a constant temperature. For example, highly specific, visualized and fast methods, such as LAMP (loop-mediated isothermal amplification), use temperatures of 60–65 °C and are based on DNA synthesis by spin-loop substitution employing DNA polymerase and two sets of primers: external primers generate a stem-loop DNA for a LAMP cycle, and internal primers are used for chain shifting during DNA amplification. The LAMP-amplified products can be detected by the naked eye, fluorescence and turbidity measurements. A turbidimeter is used to assess the turbidity of solutions as it can measure the change in turbidity due to magnesium pyrophosphate complex formation or fluorescence of amplicons upon the addition of fluorescent reagents, such as SYBR Green 1 or

**Table 3**

| Method                          | Advantages                                                                 | Disadvantages                                                                 |
|--------------------------------|---------------------------------------------------------------------------|------------------------------------------------------------------------------|
| PCR                            | Speed, cost-effectiveness, and comfort of automation                       | May lead to false detection due to similarity of *ail*, *myf*, and *yst* genes. |
| LAMP                           | Real-time detection                                                      | Requires pre-enrichment steps before PCR.                                     |
| HRMT                           | High-resolution melting technology                                         | Simple but requires sequence variants with desired melting points and labelled primers. |
| LAMP-amplified PCR             | Simple and inexpensive                                                    | Requires pre-enrichment steps before PCR.                                    |

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fluorescent metals. *Y. enterocolitica* was detected in minced pork and milk samples by using gyrB, phoP, and outU genes as the targets by the LAMP method.61,62 Zhang and co-workers employed another type of CPA, in which the testing primers were modified with biotin and FITC (fluorescein isothiocyanate) and combined with an immunoblotting assay of nucleic acid lateral flow strips. Noticeable results of detection were achieved by dipping the strips in an anti-biotin solution; two red lines appeared for positive results, whereas one red line was observed in case of false outcomes. This method is 100 times more sensitive as compared to PCR, with the detection limit of 10–100 CFU g⁻¹.61,64

Besides the development of these methods, the emergence of microarray assays brings more advancements in the detection of pathogens. In this technique, pathogens are immobilized and later detected by implementing different strategies. *Y. enterocolitica* can be detected in various samples by developing microchips with the combination of the PCR amplification of multiple genes (ail, virF, yst, and blaA) and random DNA fragments. Recently, a silicon-based optical thin-film biosensor chip was developed for the sensitive detection of 11 foodborne pathogens, including *Y. enterocolitica*. Fourier transform infrared spectroscopy (FT-IR) and matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS) are new methods developed in recent years, which can be used to identify the species and subspecies of *Y. enterocolitica*. In another study, *Y. Enterocolitica* was detected in samples due to its ability of conjugation with deferoxamine by the cell surface siderophore receptor of bacterial cells. In this method, deferoxamine-modified bovine serum albumin was fixed on gold plates and upon introduction of bacterial cells, deferoxamine–Fe composites were formed, which were rapidly detected by dark field microscopy analysis with a detection limit of 10⁶ CFU mL⁻¹.66

Furthermore, the same phenomenon was used in another study, where silica-coated magnetic nanoparticles functionalized with feroxamine (MNP@SiO₂@NH@Fa) were fabricated; the bacterial cells were successfully captured and detected in the samples. The main problem associated with this method involves the blockage material for non-specific binding, the development of new blockage materials, and validity or updating of the method.67 A high speed, ease of use and high-throughput analysis make these techniques superior to other identification methods, which can be used to detect *Yersinia* spp. However, these methods need to be further modified to reduce their high costs and long sample preparation requirements for routine testing.

**Prevention and control**

*Y. enterocolitica* is a psychrotroph and therefore, refrigeration cannot be used to control its growth. Proper sanitation at all stages of handling and processing foods and appropriate heat treatments are essential to monitor the occurrence of foodborne yersiniosis. The consumption of raw milk or meat cooked at low temperatures should be avoided. *Y. enterocolitica*, being susceptible to heat and pasteurization, can be easily destroyed by ionizing radiation, UV-radiation, and other food preservation procedures. Recently, a pulsed light-based reduction method was introduced for pork meat: the product was kept in a benchtop sterilization system (SteriPulse-XL 3000, Model RS-3000C, Xenon Corporation, Wilmington, Massachusetts, USA) and analysed for microbial analysis. The results showed that the pulsed light treatment (20–1100 nm) significantly reduced the number of bacterial cells through the formation of pyrimidine dimers, primarily thymine dimers, and the destruction of these cells due to the photochemical effect, which inhibited cell replication and led to clonogenic death.79 In another study, virulent bacteriophages Podoviridae (*fHe-Yen3-01*) and Myoviridae (*fHe-Yen9-01, fHe-Yen9-02* and *fHe-Yen9-03*) were applied to raw pork samples to control *Y. enterocolitica* contamination. The results showed that *fHe-Yen9-01* was the most effective bacteriophage with the maximum host range.80 *Y. enterocolitica* biotype 4 has the ability of biofilm formation in meat processing units; sodium hypochlorite and quaternary ammonium compounds (QAC) were found to be the most active compounds to avoid these biofilm formations.81

*Yersinia* produces two types of β-lactamases (enzymes that hydrolyse the β-lactam ring of β-lactam antibiotics) and is thus resistant to the penicillin group of antibiotics. Due to β-lactamase activity, this pathogen shows natural resistance to penicillin but is considered sensitive to newer antimicrobial β-lactam antibiotics such as ceftriaxone, cefazidime, and moxalactam. Moreover, the *Y. enterocolitica* strains are the most susceptible to aminoglycosides, extended-spectrum cephalosporins, trimethoprim-sulfamethoxazole, tetracyclines, and fluoroquinolones.4,15,71 *Y. enterocolitica* is equally sensitive to imipenem and aztreonam antibiotics. Broad-spectrum cephalosporins are also effective against extraintestinal infections. Resistance to fluoroquinolones occurs sporadically either due to a mutation in the gyrA gene or efflux mechanisms. Multi-resistant strains have been reported among pig strains belonging to bioserotype 4:O3.82 The inactivation of multidrug-resistant *Y. enterocolitica* on a stainless steel surface by cold atmospheric pressure plasma (CAP) treatment with peak-to-peak voltage of 10 kV and sinusoidal waveform of 2 kHz was also reported, where the direct antigen detection provided rapid and specific identification results. However, the specificity of these methods is limited due to the shelflife, batch to batch alteration and quality of antibodies. Propidium iodide and SYTO 9 were also used as staining and cell destruction agents according to some reports. The significant duration-dependent reduction ranged from 1.68 ± 0.17 to 2.80 ± 0.17 log.84 Still, more investigations and the development of new strategies are required for the control and prevention of *Y. enterocolitica*.

**Concluding remarks**

Yersiniosis is a gastrointestinal ailment triggered by *Y. enterocolitica*. At present, many cultural, immunological, and molecular techniques are being used for the detection of this pathogen. The use of CIC and SSDC agar is dominant for its
isolation and identification, but carrying out these methods is cumbersome, lengthy and unreliable and needs biochemical confirmation. Therefore, the detection of Y. enterocolitica in food was improved by introducing serological and molecular methods such as IMS, bacterial hybridization, PCR, microarray assays, and LAMP. The low incidence of Y. enterocolitica in food is related to the insensitivity of the current detection methods. In the future, approaches based on chromogenic/fluorogenic media, immuno-PCR, biosensors, flow cytometry, nucleic acid sequencing and aptamer-based detection can be optimized and developed for more accurate and routine diagnostics of Y. enterocolitica. These will serve as powerful tools for authorities to accurately estimate and reduce the incidence of Yersiniosis.

Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| SNPs         | Single nucleotide polymorphism |
| CPA          | Cross priming amplification |
| LAMP         | Loop-mediated isothermal amplification |
| CDC          | Centre for disease control and prevention |
| QAC          | Quaternary ammonium compound |
| IMS          | Immunomagnetic separation |

Conflicts of interest

There are no conflicts to declare.

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