Emerging Pathogens in Meat and Poultry

U.S. must step up efforts to rapidly detect and control new foodborne hazards
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Any opinions and conclusions expressed herein are those of The Pew Charitable Trusts and do not necessarily represent the views of the above individuals.

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Overview

Meat and poultry are among the leading vehicles for foodborne illnesses around the world and are responsible for sickening more than 2 million Americans each year. The pathogens that cause these infections are typically zoonotic (meaning they can be transmitted between animals and humans) and can be introduced at any point along the food chain—from when the animal is raised, to the day of slaughter and beyond, up to the moment the meat or poultry product is consumed.

A significant number of pathogens can be transmitted to humans through meat and poultry, and the risks have changed over time. The public health threat posed by some pathogens has diminished, while others have persisted for decades. New, often more virulent strains of existing disease agents continue to emerge, along with previously unknown pathogens such as the Middle East respiratory syndrome coronavirus (MERS-CoV). When such truly new pathogens emerge in one part of the world, many questions remain initially unanswered, including how they are transmitted, which animal species they can infect, whether they may extend their geographic range, and whether transmission through meat and poultry may be possible. MERS-CoV, for instance, is currently emerging in camels and humans in the Middle East; what, if any, risk this virus may pose to the U.S. meat supply going forward is difficult to predict. Some emerging pathogens, such as E. coli O157:H7, have eventually developed into major food safety concerns. For others, such as the hepatitis E virus (HEV), the importance of exposure through meat is still uncertain. Given the dynamic properties of foodborne pathogens, one thing appears certain: New foodborne risks inevitably will develop.

Ensuring the continued safety of America’s meat and poultry supply requires an agile, adaptable system that is able to detect, assess, and control both emerging and established risks. Because the pathogens that may emerge in the future cannot be known today, that is the central challenge to building a strong food safety oversight system.

This study reviews microbial hazards and risks in the U.S. meat and poultry supply that have emerged, are emerging, or that evidence suggests may emerge in the future. The study’s goals are to identify factors that favor the occurrence of emerging pathogens (EPs) and pinpoint traits that EPs transmitted through meat and poultry may share; characterize the challenges these pose, be they scientific, technological, or regulatory; and determine mechanisms that might facilitate the expeditious detection, characterization, and control of such EPs.

For the purposes of this report, EPs are defined as new microbial hazards to which significant exposure to the public through meat or poultry is possible or likely, known hazards to which new or increased exposure is possible or likely, or known hazards to which human susceptibility is increasing.¹ Unlike other definitions of EPs,² this one includes pathogens such as Salmonella that have not increased in overall occurrence but have strains with new traits that continue to emerge.

The history of how microbial hazards have emerged as foodborne pathogens provides valuable lessons learned for the study of future disease emergence. For this reason, this report includes pathogens, such as Listeria monocytogenes or Yersinia enterocolitica, that have become widely established in the meat and poultry supply and are no longer typically thought of as emerging. Studying the history of their emergence provides useful insights about the detection, characterization, and control of EPs. The report also includes EPs that have the potential to emerge in meat and poultry in the future—because they have recently emerged in other parts of the world, for instance—even if questions remain about transmission through these food products. The EPs discussed in the report can be broadly divided into six categories:
• New, previously unknown pathogens.
• Evolving strains of established pathogens.
• Known pathogens with a stronger transmission component through meat and poultry than previously understood.
• Foodborne pathogens affecting susceptible subgroups of the population that are growing in size.
• Previously unknown pathogens with suspected, but not yet established, transmission through meat and poultry.
• Pathogens common in other parts of the world that may present a future emergence threat in the U.S. meat and poultry supply.

These categories are more fully explained in the Background section.

Because many EPs have traits in common, this report begins with a general discussion of EPs, primarily those associated with meat and poultry, and then focuses on issues resulting from the emergence of five major foodborne pathogens for which considerable data are available: *Campylobacter*, pathogenic *E. coli* (Shiga toxin-producing *E. coli* [STEC] as well as antimicrobial-resistant non-STEC *E. coli*), *Listeria monocytogenes*, *Salmonella*, and *Toxoplasma gondii*. The report discusses the potential effects of changes in population susceptibility and evolving strains for each pathogen, including the development of any antimicrobial resistance.

The report also explores a number of EPs known or believed to be hazards associated with the consumption or consumer handling of meat and poultry products, as well as EPs that do not occur in the U.S. today but that may extend in range and therefore pose a potential threat in the future.

The focus of this report is EPs in meat and poultry products derived from major (i.e., cattle, swine, chicken, and turkey) as well as minor (e.g., sheep, goat, ostrich, and duck) food-producing species. However, the report is not limited to these. To assess the potential impact of new EPs on all aspects of society, and to gain insights into the effectiveness of EP response, selected examples beyond the scope of meat and poultry are discussed, where appropriate, including the 1998 Nipah virus outbreak that occurred in Malaysia, the 2003 SARS–coronavirus epidemic, and the 2009 H1N1 pandemic. A discussion of the Zika virus outbreak is not included in this report because, at the time of publication, the Zika epidemic was still ongoing. When the epidemic ends, the analysis is likely to hold valuable lessons for the study of disease emergence and pandemic preparedness and response. This study ends with recommendations for implementing a more holistic approach to food safety, one that acknowledges the dynamic and interconnected ecology of microorganisms, humans, food animals, and wildlife and their respective environments.

These recommendations fall into four categories. All are explored in detail at the report’s conclusion:

• **Prediction.** Support efforts to understand what factors lead to the emergence of new pathogens and how the government and stakeholders have responded to such pathogens in the past. This will provide valuable insights for predicting and mitigating the risk of future disease emergence. It includes supporting research to uncover weaknesses in current food production practices and identifying and monitoring trends that may lead to disease emergence in the food chain and beyond.

• **Detection.** Build or enhance surveillance systems and diagnostic tools that are able to detect EPs early and that can reliably distinguish them from other microbes that do not pose a public health risk. This includes collaborating with veterinarians, the food industry, and academia to improve detection and surveillance.

• **Capacity building.** Develop agile regulatory approaches, tools, and infrastructure to foster quick EP responses
in the face of uncertainty. This includes building necessary relationships before a new outbreak occurs, so stakeholders can collaborate quickly and efficiently, along with improving coordination of emerging disease preparedness efforts among local, state, federal, and international partners.

- **Leadership and oversight.** Determine where responsibility lies for the oversight of emerging disease preparedness activities and how such efforts will be evaluated. This includes identifying strategies for reconciling competing and possibly conflicting priorities such as the allocation of scarce resources between preparing for low-probability/high-risk emergence events and monitoring common occurrences such as emerging multidrug-resistant strains.

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**Scope and Focus of This Report**

This report discusses microbial hazards associated with meat and poultry consumption. While many of the pathogens described in the report can also be transmitted through other food vehicles (e.g., contaminated produce, raw milk, or undercooked eggs), the report focuses on direct human health risks associated with the handling or consumption of meat and poultry products in the United States. Excluded from this report are issues exclusively associated with seafood, produce, game meats, eggs, and other foods, as well as concerns primarily related to imported foods or agricultural production outside the United States. Environmental impacts not directly affecting human health are also excluded. However, because of data scarcity and knowledge gaps for many newly emerged pathogens, we chose to discuss selected relevant observations associated with related food vehicles beyond meat and poultry (e.g., milk and dairy products) that are informative for evaluating the public health risks associated with EPs in meat and poultry. In addition, selected non-foodborne epidemics are discussed as they illustrate the impact of new EPs on society and the effectiveness of rapid pandemic response.

The report does not discuss the impact of climate change on EPs—an overarching, complex topic that warrants its own discussion, which is well underway elsewhere. For similar reasons, the report limits its discussion of antimicrobial resistance to the direct health risks associated with resistant pathogens entering the food supply; indirect effects, such as the potential sharing of resistance genes with environmental pathogens, are not discussed. Emerging animal diseases such as porcine epidemic diarrhea (PED) that do not pose a human health risk are also beyond the scope of the report and therefore not discussed here.

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**Background**

Meat and poultry are among the leading vehicles for foodborne illnesses in the United States. A study focusing on 14 of the most important foodborne pathogens in the U.S. found that beef, pork, and poultry products are responsible for more than 2 million Americans getting sick each year, with an annual cost exceeding $5.7 billion. The U.S. Centers for Disease Control and Prevention (CDC) has found that meat and poultry commodities account for 40 percent of bacterial foodborne illnesses. Specific public health risks associated with meat and
poultry have changed over time, however. This report focuses on emerging and potentially emerging microbial hazards to human health that are associated with meat and poultry produced in the United States.

For the purpose of this report, hazards are biological agents (i.e., bacteria, viruses, parasites, and prions) that are reasonably likely to cause harm if present in meat or poultry destined for human consumption. By contrast, risk is defined as the probability that exposure to a hazard will lead to human harm, and therefore reflects both the hazard and the likelihood of exposure. Following the definition of emerging risks proposed by the European Food Safety Authority (EFSA), and in alignment with the definition used by the U.S. Department of Agriculture Food Safety and Inspection Service (USDA-FSIS), this report discusses pathogens that present new hazards to human health (e.g., due to the acquisition of new traits), as well as known pathogens whose importance to public health has recently surged, either because human exposure has increased or the population has become more susceptible to them. USDA-FSIS defines an emerging food safety risk as “resulting from a newly identified hazard to which significant exposure may occur or from an unexpected new or increased significant exposure and/or susceptibility to a known hazard.”

Notably, while some of these EPs are relatively well documented, the public health risks from other pathogens remain unclear, and new risks and hazards are likely to continue to emerge.

We have included foodborne pathogens that have emerged since 1970 because the history of their emergence is illustrative of the challenges associated with the detection, characterization, and control of EPs. As a result, this report classifies certain well-established pathogens, such as *Listeria monocytogenes* or *Yersinia enterocolitica*, as emerging. Concerns about increases in population susceptibility or exposure may eventually lead to a recharacterization of these well-established pathogens as emerging.

While microbial hazards in meat and poultry today appear similar to those of a decade ago, there are some specific and important differences. The risks associated with meat and poultry have not remained static; while some risks have been successfully controlled or eliminated, new risks have emerged. Newly emerging hazards are often not well understood initially: A pathogen’s specific implications for food safety and public health might materialize almost immediately, as in the case of *E. coli* O157:H7, or may remain opaque for quite some time. The latter was, in fact, the case for some pathogens, such as *Listeria monocytogenes*, that are now unanimously accepted as major food safety risks.

Beef, pork, and poultry products are responsible for more than 2 million Americans getting sick each year, with an annual cost exceeding $5.7 billion.
The emergence of new foodborne risks can, for instance, be driven by the acquisition of new virulence or resistance characteristics, by demographic changes and associated increases in population susceptibility, or by the emergence of pathogens in a new host species or geographic region. In addition, new scientific knowledge may lead to the re-evaluation and re-prioritization of known risks.

This report discusses a number of EPs specific to the U.S. meat and poultry supply. The EPs can be broadly divided into six categories, with distinct risk factors and opportunities for lessons learned:

1. Pathogens such as *Yersinia enterocolitica*, bovine spongiform encephalopathy (BSE), and *Cryptosporidium*, which have emerged as food safety concerns since the 1970s, for which the proliferation is documented in contemporaneous reports, and the epidemiology has been studied fairly extensively;

2. Evolving strains of established foodborne pathogens—such as *Campylobacter* species or multidrug-resistant non-O157 STEC and *Salmonella* strains—that have recently emerged and are likely to continue presenting new food safety challenges;

3. Known pathogens such as *Toxoplasma gondii*, about which modern research suggests a potentially stronger foodborne transmission component, or greater danger to public health, than was previously understood;

4. Pathogens such as *Listeria monocytogenes* that primarily affect highly susceptible population subgroups. Their importance may be increasing as susceptibility expands in the U.S. population due to an aging population or other demographic factors, or as the importance of new food vehicles and exposures is being recognized;

5. EPs whose transmission through meat and poultry may be possible but has not yet been firmly established, such as *Arcobacter butzleri*, *Clostridium difficile*, *Helicobacter pylori*, *Mycobacterium paratuberculosis*, multidrug-resistant *Staphylococcus aureus*, and hepatitis E virus; and

6. Pathogens not currently found in the U.S. that are emerging in other parts of the world and that may be transmissible through meat or poultry, and therefore potentially pose a future threat to the U.S.; these include human-pathogenic avian influenza virus, Crimean-Congo hemorrhagic fever virus, MERS-CoV, and Rift Valley fever virus.

The public health risks associated with meat and poultry products, science’s understanding of their causes and human health impacts, and the various regulatory responses to them have changed considerably over time. In A.D. 900, Byzantium’s Emperor Leo VI forbade—under threat of exile and loss of all possessions—the making and eating of blood sausage, prepared in pigs’ stomachs and then smoked. It was believed that miasmas, or infectious vapors, emitted from the sausages would result in a paralytic infection. This disease was called “botulism,” after “botulus,” the Latin word for sausage. Botulism’s true cause, a toxin produced by a bacterium named *Clostridium botulinum*, would not be discovered until 1895, when Emile Pierre van Ermengem at the University of Ghent in Belgium studied an outbreak associated with smoked ham.

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The public health risks associated with meat and poultry products, science’s understanding of their causes and human health impacts, and the various regulatory responses to them have changed considerably over time.
Figure 1
How and Why Pathogens Emerge
*E. coli* O157:H7 and ground beef in the U.S.

The population becomes **more susceptible** to an existing pathogen, and more people fall ill.

An existing pathogen acquires **new traits** and becomes a greater threat to public health.

A pathogen suddenly appears in a **new host**—a pathogen that infected one species now infects a new species.

A pathogen emerges in a **new geographic region** that is not equipped to deal with it.

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**E. coli O157:H7 and Ground Beef in the U.S.**

There is perhaps no better example of an emerging foodborne pathogen than *E. coli* O157:H7, also known as Shiga toxin-producing *E. coli*, or STEC.

**Emergence**

- In 1975, *E. coli* O157:H7 was isolated from a patient but was not recognized as a foodborne pathogen until seven years later.

- In 1982, the Centers for Disease Control and Prevention (CDC) identified *E. coli* O157:H7 as a new foodborne pathogen after it caused two outbreaks of infections associated with ground beef hamburgers. Illnesses were increasingly detected over the next few years.

- In January 1993, four children died and more than 500 people were sickened after eating undercooked contaminated hamburgers from Jack in the Box restaurants.

**Government response**

- In 1993, the Food and Drug Administration (FDA) revised the Model Food Code for restaurants with new cooking temperature guidelines for ground beef.

- In 1994, *E. coli* O157:H7 became a nationally notifiable disease—in other words, collecting regular information on *E. coli* infections was considered necessary to protect public health.

- Also in 1994, the U.S. Department of Agriculture (USDA) declared *E. coli* O157:H7 in raw ground beef an “adulterant” (i.e., contamination renders ground beef unfit for human consumption).
In 1996, USDA mandated prevention-based regulations for meat and poultry plants known as the Pathogen Reduction/Hazard Analysis and Critical Control Point (PR/HACCP) rule.†

In October 1994, USDA’s Food Safety and Inspection Service began a microbial testing program in raw ground beef; this reduced the risk of contaminated product reaching the consumer.†

Since 1997, CDC has included E. coli O157:H7 in its foodborne pathogen fingerprinting database, and has improved surveillance of outbreaks.‡

The National Livestock and Meat Board developed objective measures for the “doneness” of hamburgers and encouraged automated cooking systems.‡

Industry implemented control steps, such as carcass rinses and steam vacuum systems, to minimize the risk of E. coli O157:H7 contamination during slaughter.§

Beef industry response

The National Livestock and Meat Board developed objective measures for the “doneness” of hamburgers and encouraged automated cooking systems.‡

Industry implemented control steps, such as carcass rinses and steam vacuum systems, to minimize the risk of E. coli O157:H7 contamination during slaughter.§

All of the measures together led to an estimated 50 percent decrease in E. coli O157:H7 infections from 1997 to 2011.||
This report distinguishes the following four broad categories of human foodborne exposure:

- **Direct foodborne exposure**: from consuming or handling raw or undercooked beef, pork, or poultry products.
- **Direct non-foodborne exposure**: through contact with or proximity to live animals or their environments.
- **Indirect foodborne exposure**: through produce contaminated with manure via environmental pathways (e.g., wind, water) or farming practices (e.g., application of manure to crop fields).
- **Indirect non-foodborne exposure**: via interactions with natural environments (water, soil, or wild animals) polluted by nearby animal production facilities.

It is very difficult to characterize the overall and relative importance of these pathways. Due to limitations in data and knowledge, this report focuses on direct foodborne exposure, although the report does touch on other exposure routes where necessary.

Emerging infections have commonly been defined as having newly appeared in a population or having existed but are rapidly increasing in incidence or geographic range. A broader definition includes a new hazard to which significant exposure is possible or likely, a known hazard for which new or increased exposure is possible.
or likely, or a known hazard to which human susceptibility is increasing.\textsuperscript{15} Included in this broader definition are pathogens, such as \textit{Salmonella}, for which the overall prevalence has remained constant but strains with new properties periodically emerge. Exactly how long a hazard remains “emerging” may differ and is subject to varying interpretations. Some EPs may eventually become endemic and will therefore no longer be emerging, while others will be effectively eradicated or controlled.

Because EPs present a significant economic and public health burden, considerable efforts have been dedicated to studying them in an effort to better prepare for them and ultimately predict when they will appear. Most EPs share certain traits that facilitate their emergence.\textsuperscript{16} Additionally, scientists have correlated socio-economic, environmental, and ecological factors with the occurrence of EPs.\textsuperscript{17} The complex interactions among these factors, and their roles in facilitating emergence events, are easiest understood in the context of the epidemiologic triangle, also referred to as the host/agent/environment model of disease causation. Disease occurs when an agent (i.e., pathogen) meets a vulnerable host (i.e., human) in an environment that facilitates disease transmission (e.g., cross-contamination of meat in a retail delicatessen). (See Figure 2.)

\textbf{Figure 2}
\textit{Epidemiologic Triangle}

\begin{center}
\begin{tikzpicture}
\node (host) at (0,0) {Host};
\node (agent) at (-2,-4) {Agent};
\node (environment) at (2,-4) {Environment};
\draw[->] (host) -- (agent);
\draw[->] (host) -- (environment);
\draw[->] (agent) -- (environment);
\end{tikzpicture}
\end{center}


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Disease emergence typically results from changes in the intricate interplay between host, agent, and environment.\textsuperscript{18} However, different factors may be associated with emergence, depending upon whether it is due to a change in host range (e.g., an animal disease becomes capable of infecting humans), the acquisition of new traits without a corresponding change in host range (e.g., acquiring of antimicrobial resistance or virulence traits), or its emergence in a new geographic area.\textsuperscript{19}
Traits associated with the pathogen (agent)

Microorganisms are constantly evolving. Changes in genetic structure may make certain strains of a pathogen more efficient and better able to survive within changing environments or may allow them to survive in new environments. Pathogens can evolve into novel strains, become more virulent, or acquire resistance to antimicrobial drugs or other treatments. New pathogens may be discovered, or existing pathogens may acquire the ability to infect new hosts. Similarly, a pathogen that was previously controlled or had naturally declined can re-emerge or be reintroduced.

Certain microbial characteristics are thought to increase the risk of emergence. Zoonotic pathogens (i.e., those that can be transmitted between animals and humans) have been responsible for approximately 60 percent of infectious diseases emerging in a new host species since 1940 and are thought to pose a particularly high risk. Likewise, microbes that can infect multiple species, evolve at an inherently high mutational rate (e.g., RNA viruses), or are predisposed to the acquisition of genetic material (e.g., through horizontal gene transfer or reassortment) are thought to be at increased risk of emergence events.

A number of foodborne EPs have surfaced over the past 40 years, and in recent years novel strains of them, as well as new pathogens that potentially could be transmitted through meat and poultry, continue to be discovered. Serotypes (i.e., specific subtypes of the bacterium defined by serological measures) associated with human illness have shifted over time, new virulent subtypes have emerged, and many strains have acquired antimicrobial-resistance genes.

Traits associated with the exposed individual or population (host)

How easily a pathogen can invade a human or animal is thought to be an important factor in disease emergence. A variety of risk factors have been identified, both on the individual level (e.g., age, underlying disease) and across populations (e.g., population structure, community composition, resilience toward invasion). Individuals may, for example, differ in their susceptibility to infection because of their immunological history (e.g., vaccination, prior exposure), genetic predisposition, exposure to pharmaceutical drugs (e.g., antacids, immune-modulating drugs), general health status (e.g., obesity, nutritional health), or because they have been affected by other diseases or conditions (i.e., “comorbidities” such as HIV/AIDS or diabetes mellitus).

Susceptibility to infection with foodborne pathogens increases with older age, diabetes mellitus and other comorbidities, as well as conditions such as obesity that lead to decreased immune responses. In the U.S., the number of people aged 65 and older is expected to increase 135 percent between 2000 and 2050, with the subpopulation of people aged 85 and older increasing almost 350 percent. Additionally, more than one-third of Americans are obese, and between 1990 and 2008 the rate of diabetes doubled.

Many foodborne pathogens disproportionately affect the young, old, pregnant, and immunocompromised. In the future, demographic shifts resulting from population movements or differences in birth rates, as well as changes in the prevalence of underlying diseases or conditions, may lead to shifts in the susceptibility to foodborne infections.

Traits associated with the environment

A variety of environmental factors have been linked to disease emergence, including climatic factors (e.g., temperature, rainfall, and humidity), seasonality, and the presence of environmental barriers such as rivers or mountain ranges. Environmental factors may, for instance, prolong pathogen survival in the environment.
or contribute to a pathogen's dispersal through wind or rain runoff. The environment can also affect the seasonal presence of vectors such as ticks and mosquitoes, reduce host resistance (e.g., through heat stress or malnutrition), or lead to increased contact rates among people and/or animals (e.g., crowding during the winter). Therefore the contributions of host, pathogen, and environment to disease emergence are intricately linked.

A foodborne pathogen’s “environment” consists of the complete exposure pathway from its introduction into the food supply, to its eventual consumption by people. Increased exposure to microbial hazards can result from changes in food production or consumption, which can open new avenues for the introduction, proliferation, and transmission of pathogens.

Changes in food production and consumption patterns may lead to complex changes in disease epidemiology, even if these changes may not always be directly apparent from an analysis of the overall disease burden. For example, even if the overall incidence of a certain foodborne disease is not changing, the risks associated with one specific food commodity may be emerging. If the risks associated with other food commodities are simultaneously becoming better controlled, this can mask increases in disease incidence caused by the emerging risk. Demographic shifts can result in changes in food preparation and consumption and may affect associated risks. To better understand and be able to quickly react to an emerging foodborne risk, it is imperative for scientists to analyze emerging trends, in both overall pathogen incidence and in food vehicles associated with particular pathogens. It is equally important to have reliable methods for linking foodborne illnesses back to food vehicles—an often challenging task—and to understand the root causes that ultimately came together to cause the illnesses.

It is important to note that in some cases the increased recognition of an established hazard—for example, as a result of improved methods for pathogen detection or characterization, or through expanded surveillance—may be difficult to distinguish from the emergence of a new hazard. In fact, in many cases evidence suggests that pathogens that emerge in a new host species require time to reach full adaptation and often circulate for months or years before being recognized.28

Another point worth clarifying is that many EPs, by their very nature, are not as well understood as long-standing problems. Scientific evidence may be scarce, and expert communities may be split in their assessments of risk. For instance, the zoonotic potential and host range of EPs (i.e., the ability of a new animal disease to infect humans, and the risk of foodborne transmission) are often uncertain for newly emerged pathogens.

New and emerging foodborne pathogens

Foodborne diseases are a significant public health challenge in the United States. The CDC estimates that microorganisms transmitted through food sicken one in six Americans each year, resulting in about 128,000 hospitalizations and 3,000 deaths.29

Of the 9.4 million cases of foodborne illness that the CDC ascribes to 31 major pathogens, over 90 percent are due to only five—norovirus, Salmonella, Clostridium perfringens, Campylobacter, and Staphylococcus aureus. (See Figure 3.)30 Similarly, five pathogens (Salmonella, norovirus, Campylobacter, Toxoplasma gondii, and Shiga toxin-producing E. coli) are estimated to cause 88 percent of hospitalizations due to these pathogens. (See Figure 4.) And five pathogens (Salmonella, T. gondii, Listeria monocytogenes, norovirus, and Campylobacter) are estimated to cause 88 percent of deaths attributable to these pathogens. (See Figure 5.) Beef, pork, poultry, and other animal products are estimated to be among the most important food vehicles for all of these pathogens, with the exception of norovirus and Staphylococcus aureus.31
Figure 3
5 Pathogens Cause the Majority of Foodborne Illnesses in the U.S.
Estimated share of U.S. foodborne illnesses caused each year by 31 major pathogens

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Figure 4
5 Pathogens Cause the Majority of Hospitalizations Due to Foodborne Illness in the U.S.
Estimated share of U.S. hospitalizations caused each year by 31 major pathogens

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Many of the most common foodborne pathogens in the U.S. today were discovered or associated with foodborne exposure in the past 50 years. This fact is likely due at least in part to advances in microbiological and diagnostic science and disease surveillance systems. Examples of major pathogens associated with meat and poultry consumption include *E. coli* O157:H7 and other Shiga toxin-producing *E. coli*, *Campylobacter jejuni*, and *Listeria monocytogenes*. (See Table 1.) Disease surveillance data, available from the CDC’s Foodborne Diseases Active Surveillance Network (FoodNet) since 1996, indicate an intricate picture of the dynamics of foodborne infectious diseases. For many of the major known foodborne pathogens that are often transmitted through meat and poultry exposure, incidence initially declined in the late 1990s and early 2000s (possibly related to the USDA’s implementation of the PR/HACCP rule in 1996) and has since plateaued or slightly increased. Importantly, surveillance data are vulnerable to external influences, such as changes in resource availability, and therefore have to be interpreted carefully. In addition, changes in industry practices can affect surveillance trends. The *E. coli* O157:H7 outbreak linked to ground beef served in Jack in the Box restaurants, for example, led to changes in government regulation and industry practices, increased surveillance for the pathogen, and a decrease in the burden of STEC cases.

Figure 6 shows trends in the incidence of the major foodborne pathogens in the U.S., which are often transmitted through meat and poultry consumption.
Figure 6
Trends in Incidence of Major Foodborne Pathogens
Relative rates of laboratory-confirmed infections in the U.S. per year, compared with rates for 1996-98

New serotypes and molecular subtypes of common foodborne pathogens have emerged in recent years and continue to do so, with growing concerns about the impact on public health and importance of these strains. Examples include non-O157:H7 STEC and *Campylobacter* strains other than *C. jejuni*. In addition, pathogens have emerged with resistance to one or more antimicrobial drugs. These have been responsible for a number of notable foodborne outbreaks and reflect another emerging public health concern.

A variety of pathogens are newly emerging in the U.S. or around the world. Others have only recently become a concern for meat and poultry exposure, either from ingestion (i.e., oral exposure) or other food-associated exposure pathways—for instance, dermal contact between consumers and the contaminated meat they handle.

Which pathogens should be considered as emerging or potentially emerging hazards in the U.S. meat and poultry supply is somewhat subjective. The pathogens included in this report were selected because they were identified in one of the source documents in Table 1, or because they were identified as important potential hazards by two subject matter experts involved in writing this report based on their interpretation of the scientific literature (see box for more details).

Table 1 shows emerging hazards of potential concern with exposure to meat or poultry products. As described in the overview, these include pathogens present in the U.S. as well as ones that may pose a risk if they are introduced to the U.S. at a future time. The hazards entail foodborne pathogens as well as pathogens such as Methicillin-resistant *Staphylococcus aureus* (MRSA) that may pose a food-handling risk.
Pathogens that pose clear, well-understood foodborne risks associated with meat and poultry consumption are further discussed in the section starting on page 17. They are the primary focus of this report.

Pathogens for which foodborne transmission with meat and poultry products remains subject to debate, that have scarce transmission data, or are not endemic in the U.S are discussed in the section starting on page 38.

Table 1
Emerging Pathogens Potentially Transmitted Through Exposure to Meat or Poultry

<table>
<thead>
<tr>
<th>Reason for Inclusion</th>
<th>Meat and poultry associated</th>
<th></th>
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<tbody>
<tr>
<td><strong>Bacterial pathogens</strong></td>
<td></td>
<td></td>
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<tr>
<td>Arcobacter butzleri</td>
<td></td>
<td></td>
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<tr>
<td>Campylobacter jejuni, C. Fetus</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E. coli (STEC &amp; non-STEC)</td>
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<tr>
<td>Listeria monocytogenes</td>
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<tr>
<td>Yersinia enterocolitica, pseudotuberculosis</td>
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<tr>
<td><strong>Viral pathogens</strong></td>
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<tr>
<td>Hepatitis E virus</td>
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<tr>
<td>BSE/vCJD/TSE*</td>
<td></td>
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<tr>
<td>Avian influenza virus (AIV)*</td>
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<tr>
<td>Crimean-Congo hemorrhagic fever virus (CCHF)*</td>
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<tr>
<td>Hepatitis E virus</td>
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<tr>
<td>MERS-CoV*</td>
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<tr>
<td>Rift Valley fever virus (RVFV)*</td>
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<td><strong>Parasites and prions</strong></td>
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<tr>
<td>Cryptosporidium</td>
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<tr>
<td>Toxoplasma gondii</td>
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<td></td>
</tr>
<tr>
<td>BSE/vCJD/TSE*</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Not currently a risk in the U.S., but future introduction is possible.
† Transmission through a route other than foodborne (e.g., dermal).

Note: Bold typeface indicates pathogens that pose a clear risk with meat or poultry; light typeface indicates pathogens for which transmission with meat and poultry products remains subject to debate or that have scarce transmission data or are not currently endemic in the U.S.
Methodology Used to Develop Table 1

A review of the scientific literature did not generate one single, comprehensive list of pathogens of emerging concern for the U.S. meat and poultry supply, defined for the purpose of this study as products derived from major (i.e., cattle, swine, chicken, and turkey) as well as minor (e.g., sheep, goat, ostrich, and duck) food-producing species. However, two relevant lists were identified in scientific publications. (Pathogens that appeared on both lists were included twice in Table 1.) The first, falling under reason for inclusion #1 (based on Emerging Foodborne Pathogens and Problems: Expanding Prevention Efforts Before Slaughter or Harvest†), lists pathogens that have emerged as foodborne risks since the 1970s and is based on a literature review. The second, identified by reason for inclusion #2 (based on “Drivers of Emerging Risks and Their Interactions in the Domain of Biological Risks to Animal, Plant and Public Health: A Pilot Study”††), highlights emerging biological risks in the European Union, based on the opinion of two expert panels (the Biological Hazard and Animal Health and Welfare panels) convened by the European Food Safety Authority to advise on scientific issues relevant to food safety and foodborne diseases in the European Union. Emerging issues considered in this opinion include new pathogens introduced or reintroduced in the EU, pathogens with possibly increased exposure, and pathogens identified because of changed susceptibility in the population.

We included only those pathogens from the two lists that may be of concern to the U.S. meat and poultry supply, using the following criteria for exclusion: those that cannot be transmitted to humans through meat or poultry (e.g., because they can infect a limited range of animal species such as fish, are animal but not human pathogens, or are known to be transmitted only through routes other than food); pathogens associated with game meat or certain nonconventional production systems that have decreased in incidence in the U.S. in recent decades (e.g., Trichinella spiralis); and pathogens not present in Europe, but commonly occurring in the U.S. (e.g., Corynebacterium paratuberculosis), because they are an established, rather than emerging, risk in the U.S. General risks identified in the two lists but not associated with a specific pathogen (e.g., “existing viruses becoming foodborne”) were not included in the table but are discussed in the report.

The condensed list generated by this method was supplemented to capture other risks that our two experts who contributed to this study consider to be of emerging or potentially emerging concern in the U.S. meat and poultry supply.

Major meat- and poultry-associated pathogens with emerging strains

In this section, the report focuses on the most common pathogens associated with meat and poultry products: *Campylobacter*, pathogenic *E. coli*, *Listeria monocytogenes*, *Salmonella*, and *Toxoplasma gondii*. The report discusses the risks these pathogens pose to people today, as well as emergence factors that may define how science will address them tomorrow. Because the threat and public health relevance of STECs differ from that of antimicrobial-resistant non-STEC *E. coli*, we discuss these two separately. Notably, some of these pathogens have emerged since 1970 but have now become well-established food safety risks in the U.S. Because of their value for lessons learned, we have included these hazards in the report even though they may no longer be typically considered EPs.

These pathogens were chosen for one of two reasons (see Table 1 and the corresponding methodology text box for a full rationale of inclusion criteria):

**Reason #1:** They have been recognized as foodborne pathogens only since 1970 and may therefore contain valuable lessons learned.

**Reason #2:** Expert panels convened by the European Food Safety Authority identified them as emerging or re-emerging diseases. (A comparable expert panel convened by U.S. authorities was not available at the time the report was written.)

*Campylobacter*

**Reason for inclusion: #1**

Although infections with *Campylobacter* species have probably caused human illness for centuries, it was not until 1968 that this bacterium was first isolated from stool samples of patients with diarrhea, and *Campylobacter jejuni* was first identified as a human diarrheal pathogen in 1973. The study of campylobacteriosis was made possible in large part by diagnostic advancements. Selective growth media and controlled growth conditions (e.g., under controlled gas atmospheres) were developed and increasingly used in the 1970s; by the late 1980s, *Campylobacter* was recognized as the leading cause of bacterial gastroenteritis in the world. In the United States, *C. jejuni* remained the leading cause of bacterial foodborne illness until about 2001, when it was surpassed by *Salmonella*.

Recent changes in the epidemiology of foodborne campylobacteriosis were brought about by emerging strains beyond *C. jejuni* and *Campylobacter coli*. In addition, new data indicate that the use of certain prescription drugs may lead to an increased disease risk. New scientific knowledge has also led to a reconsideration of the most important transmission pathways.

**Disease burden**

The cost of foodborne campylobacteriosis is estimated at over $1.9 billion each year in the United States alone, from an estimated 850,000 cases of illness, over 8,000 hospitalizations, and nearly 80 deaths. In a 2012 ranking of major foodborne pathogens across all food commodities, *Campylobacter* in poultry was found to cause a greater disease burden than any other pathogen/food combination, being responsible for an estimated $1.3 billion in costs of illness annually. One reason for the high cost associated with *Campylobacter* infection is the
risk that after infection, patients can develop Guillain-Barré syndrome, an autoimmune disorder that leads to temporary paralysis and may cause permanent nerve damage.44

*Campylobacter* control is complicated by the low infectious dose (i.e., low number of bacteria needed to cause infection) and the resulting high risk of cross-contamination during and after slaughter (e.g., through shared cutting boards). However, the U.S. rate of campylobacteriosis dropped by half at the beginning of the 21st century, coinciding with and likely resulting, at least in part, from new practices implemented in poultry processing operations, such as the chlorination of water baths and chiller tanks, and improved sanitation during slaughtering.45 In 1997, the reported incidence peaked at 24.6 illnesses per 100,000 people per year, and subsequently fell to a low of 12.6 per 100,000 in 2003. As shown in Figure 7, the incidence has remained around 13 to 14 cases per 100,000 per year for more than a decade.46 Therefore, little progress has been made recently in reducing the rate of *Campylobacter* infections. Even though attributing *Campylobacter* illnesses to food vehicles has remained challenging, and raw milk is also an important food vehicle,47 these data do suggest that more needs to be done to control the *Campylobacter* risk associated with poultry.

**Figure 7**

*Trends in Culture-Confirmed Campylobacter Infections*

Incidences (per 100,000 population) of such infections in the U.S., by year

Emerging strains

Emerging Campylobacter species

While C. jejuni and C. coli are estimated to be responsible for more than 95 percent of campylobacteriosis cases, a number of emerging species have been associated with human illness. More than 10, including C. concisus, C. lari, C. upsaliensis, and C. ureolyticus, have been isolated from patients with gastroenteritis.

Many of these emerging species have complex nutrient requirements, making them difficult to grow in bacterial culture, even some culture media routinely used for C. jejuni. Thus, these strains are likely under-detected in clinical isolates and underestimated in terms of disease burden. It is possible that an as-yet-untitled species of Campylobacter may ultimately be found to be responsible for a considerable share of the large number of gastroenteritis cases without known etiology.

Furthermore, C. concisus and some other species have been recently associated with Crohn’s disease, although much more research is needed to establish causality. Exposure routes for emerging species are not well defined, but some strains have been isolated from many of the same foods as C. jejuni and C. coli.

Antimicrobial-resistant strains

Drug-resistant Campylobacter presents a considerable public health challenge and appears to be linked, at least in part, to antimicrobial drug use during food production. For example, the emergence of quinolone-resistant Campylobacter jejuni infections in the U.S. quickly followed FDA approval of fluoroquinolones for poultry use in 1995.

Minnesota public health officials documented a rise in quinolone-resistant human C. jejuni isolates from 1.3 percent in 1992 to 10.2 percent in 1998. In 1997, they were able to isolate quinolone-resistant C. jejuni from 14 percent of retail chicken samples, and matched these strains to human illnesses using molecular subtyping. The CDC began tracking quinolone-resistant C. jejuni in 1997 and by 2002 found 20 percent of clinical isolates to be quinolone-resistant.

International studies also support the role of agricultural drug use in the development of resistance: Countries with low or no fluoroquinolone usage on farms have a low prevalence of resistant infections despite the use of fluoroquinolones in human medicine, while countries with high fluoroquinolone farm usage have correspondingly higher rates of resistance in both human and animal isolates. A case-control study conducted by FoodNet found that people infected with fluoroquinolone-resistant Campylobacter were more likely than control groups to have consumed chicken or turkey.

Concerns about the direct role of the agricultural uses of antimicrobial drugs in the emergence of resistant Campylobacter led the FDA to withdraw approval for use of fluoroquinolones in drinking water for poultry effective in 2005. The bacterium’s resistance to ciprofloxacin is variable but trending down. Based on surveillance of retail chicken samples by the National Antimicrobial Resistance Monitoring System (NARMS) for enteric bacteria, rates of C. coli resistance to ciprofloxacin, one of the most commonly used fluoroquinolones in humans, dropped from nearly 30 percent in 2005 to approximately 20 percent in 2013. Ciprofloxacin resistance rates for C. jejuni in retail chicken are at an all-time low, decreasing from about 15 percent at the time of the ban to 11 percent in 2013. Rates of quinolone resistance in human isolates peaked at 26 percent in 2007 and were at 23 percent in 2013.
Changing exposure pathways

*Campylobacter* species are routinely found in cattle, pigs, sheep, and birds, with birds the most common hosts due to their higher body temperature.6 Most poultry flocks are positive for *Campylobacter*, and studies of retail poultry have found contamination rates in carcasses of up to 70 percent.62 A 2012 USDA-FSIS microbiological baseline study on chicken parts found that more than 21 percent were contaminated with *Campylobacter*.63 *Campylobacter* contamination rates in retail chicken reported by NARMS equaled 38 percent positive samples in 2013.64

Epidemiological research points to a complicated and possibly changing picture of foodborne exposure, and to vast differences in the epidemiology of sporadic infections and outbreaks. Sporadic *Campylobacter* infections seem to have distinct causes. For example, poultry is a rare cause of *Campylobacter* outbreaks, despite its importance among sporadic cases. Raw milk is the leading cause of *Campylobacter* outbreaks, and according to one CDC study, more than 60 percent of *Campylobacter* outbreaks are due to dairy products.65

By contrast, a large FoodNet case-control study of sporadic *Campylobacter* infections found only 1.5 percent of sporadic campylobacteriosis cases to be associated with unpasteurized milk.66 In the case-control study, consumption of restaurant-prepared chicken was responsible for nearly 24 percent of illnesses, and eating non-poultry restaurant meat was responsible for an additional 21 percent. Tasked with estimating the source of foodborne campylobacteriosis, a large panel of food safety experts concluded that about 8 percent of *Campylobacter* illnesses are due to dairy products; the panel assigned 72 percent of foodborne campylobacteriosis cases to poultry, 5 percent to produce, 4 percent to beef, and 4 percent to pork.67

Exposure may not be homogeneous across the United States. A number of studies have documented differences in the sources of campylobacteriosis between rural and urban populations, particularly in areas with animal production.68 A number of studies have suggested that cattle production may be a more important contributor to human campylobacteriosis than previously thought.69 Other researchers pointed to the potentially important role of flies in transmitting *Campylobacter* from areas of food animal production to local communities.70

Changing population susceptibility

The complex epidemiology of campylobacteriosis may be due, at least in part, to differences in population susceptibility. While differences in disease incidence have been clearly documented by the CDC, the geographic variability in campylobacteriosis rates in the United States could not be explained by differences in surveillance or known risk factors.71

Additionally, factors such as the increased use of therapeutic drugs such as proton pump inhibitors (PPI) among humans may lead to a future increased risk of *Campylobacter* infection. PPIs reduce gastric acid production and are prescribed for gastroesophageal reflux disease, dyspepsia, and similar conditions. A few studies in Europe have associated increased PPI use with increased rates of campylobacteriosis.72 While more data are needed, if a true association exists, greater PPI use in the United States may result in an increased risk of campylobacteriosis.

Shiga toxin-producing *E. coli* (STEC)

**Reasons for inclusion: #1 and 2**

*E. coli* is a highly diverse bacterial species. Many strains do not cause disease in either humans or animals,73 and these, referred to as “nonpathogenic” *E. coli*, are a natural part of the gut microflora.74 Many strains of pathogenic *E. coli*, however, can cause diarrheal disease or illness outside of the intestinal tract, such as urinary tract infections or meningitis.75 Among the most serious intestinal pathotypes are Shiga toxin-producing *E. coli* (STEC),
also referred to as Verocytotoxin-producing E. coli (VTEC), which cause severe gastroenteritis characterized by bloody diarrhea and vomiting. STEC infection can cause hemolytic uremic syndrome (HUS), which can result in acute kidney failure and death. HUS is also associated with long-term sequelae, including chronic kidney disease, end-stage renal disease, hypertension, and deficits to many organ systems.76

There is perhaps no better example than STEC to study the emergence of a foodborne pathogen associated with meat. In 1982, CDC investigations of two outbreaks of severe bloody diarrhea associated with ground beef hamburgers led to the identification of a strain of E. coli that produced a toxin similar to the Shiga toxin of Shigella dysenteriae as a foodborne hazard.77 Foodborne outbreaks caused by this strain—E. coli O157:H7—began to be identified with increasing frequency, but it was not until January 1993 that the pathogen received public attention in the wake of an outbreak at the fast-food chain Jack in the Box, which sickened over 700 people and killed four children.78 The USDA-FSIS subsequently declared E. coli O157:H7 an adulterant, meaning a contaminant in a product that makes it unfit for human consumption, and shortly thereafter mandated the prevention-based regulations for meat and poultry plants known as the Pathogen Reduction/Hazard Analysis and Critical Control Point (PR/HACCP) rule.

More than 100 additional STEC strains have since been identified, and these non-O157 STECs are of increasing public health concern. While many are associated with milder forms of diarrheal disease, some can cause the same severe acute and long-term sequelae as O157, including hemorrhagic colitis, hemolytic uremic syndrome, and end-stage renal disease.

### Disease burden

Foodborne E. coli O157:H7 is estimated to cause over 60,000 illnesses in the United States each year, resulting in about 2,000 hospitalizations and 20 deaths, while non-O157 STEC is estimated to cause over 100,000 foodborne illnesses annually, resulting in about 300 hospitalizations.

Foodborne E. coli O157:H7 is estimated to cause over 60,000 illnesses in the United States each year, resulting in about 2,000 hospitalizations and 20 deaths, while non-O157 STEC is estimated to cause over 100,000 foodborne illnesses annually, resulting in about 300 hospitalizations but no deaths.79 Foodborne STEC infections in the U.S. are estimated to cause approximately $300 million in economic costs.80 Worldwide, STECs are estimated to cause 2.8 million acute illnesses each year.81

CDC surveillance data show an increase in reported O157 infections from 1996 to 2000, as depicted in Figure 8, likely driven by increased reporting, followed by a decline in the early 2000s.82 Over the past decade, O157 disease incidence has remained largely unchanged.83

While the incidence of O157 disease has plateaued, reported cases of non-O157 STECs are on the rise, potentially in part because of improved
surveillance for non-O157 STECs. Figure 9 shows the sharply increasing rate of non-O157 outbreaks in the past 25 years. In another study of non-O157 STEC infections in the U.S. between 2000 and 2010, over half of the outbreaks occurred in the last four years of data.84 Within FoodNet, the number of reported isolates of non-O157 STEC equaled that of O157 in 2010. The extent to which these recorded increases reflect actual increases in disease incidence is unclear, partially because major improvements in reporting and detection, as well as an increase in testing for these strains, took place during this period.

Increases in detection and reporting capabilities were partially driven by regulatory changes. In response to an increasing incidence of infections linked to non-O157 STECs, USDA-FSIS declared the “Big Six” strains of STECs as adulterants in 2011. These are *E. coli* O26, O45, O103, O111, O121, and O145, which CDC had estimated to be responsible for more than 70 percent of non-O157 STEC infections.85 Simultaneously, the agency increased testing for these strains.

Even within these six major serogroups (i.e., serologically defined groups) of non-O157 STECs, however, the associated severity of disease varies considerably. For example, although only 16 percent (152 of 940) of all non-O157 human isolates sent to CDC in a 20-year period were found to be *E. coli* O111, this strain caused almost half (10 of 21) of the associated HUS cases.86 *E. coli* O111 is also the leading non-O157 strain associated with STEC outbreaks.87
Differences in clinical outcomes associated with non-O157 STEC infections are caused by a number of virulence factors. Of the two currently known major types of Shiga toxin (stx1 and stx2), STEC strains with the stx2 gene are associated with more severe clinical outcomes than are strains with the stx1 gene. Likewise, STEC strains expressing the virulence factor intimin (eae), which allows the pathogen to adhere to intestinal epithelial cells, have been found to cause more severe disease than strains not expressing eae. Other currently uncharacterized virulence factors may also be important in determining clinical outcomes.

Heterogeneity in strain-associated disease severity raises questions about how best to target public health interventions, including whether to focus on the serogroups most frequently isolated from human cases or on non-O157 strains most likely to cause severe disease (such as those expressing virulence genes stx2 and eae) regardless of their frequency of isolation.

Because of its magnitude, the 2011 outbreak of E. coli O104:H4 in Germany provides perhaps the strongest argument for focusing on virulence factors associated with severe clinical illness, even though the occurrence was not associated with meat or poultry. This outbreak, eventually traced back to contaminated sprouts, caused over 4,000 E. coli cases, 908 HUS cases, and 50 deaths. The novel strain that caused the outbreak was a particularly virulent hybrid of two pathogenic E. coli—a strain of an enteroaggregative E. coli (normally associated with severe diarrhea) that had acquired stx2 Shiga toxin genes from an STEC strain. It was also found to carry several antimicrobial-resistance genes, making treatment very difficult.

Figure 9
Reported Non-O157 Outbreaks Over Time

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When determining the risk of future emergence events it is important to note that horizontal gene transfers (i.e., the transfer of genetic material among bacteria in a manner other than traditional reproduction), as seen in the German *E. coli* O104:H4 outbreak, are far from uncommon. As a case in point, Shiga toxins have been found contained within the DNA of bacteriophages, or viruses that infect bacteria. Bacteriophages can excise DNA from a host bacterium and may transfer the DNA to another bacterial cell; phages may, for instance, transfer virulence genes from a pathogenic bacterial strain to a nonpathogenic strain that does not otherwise cause illness.

**Changing exposure pathways**

Beef is the leading cause of foodborne infections with O157:H7 STEC in the U.S. Farm visits and exposure to cattle have also been identified as important risk factors. In the past several years, however, there has also been increased concern about STEC contamination of leafy greens and other produce, especially after a major outbreak of *E. coli* O157:H7 infections linked to prepackaged spinach in 2006. In this case, the outbreak strain isolated from patients was matched to bags of spinach, as well as to environmental samples, feces from cattle near the spinach farm, and fecal samples from feral swine. This example emphasizes the complicated and dynamic ecology of STEC infections and the central role food animals may play in both meat and produce safety.

Although cattle, and to a lesser extent other ruminants, have been established as natural reservoirs for O157:H7 STECs, the causes of emergence are unknown. Considerable research into cattle carriage and fecal shedding of O157 has found a highly dynamic system in which prevalence within farms is extremely variable over time. Diet is also important; the type of grain and processing method, forage quality, and use of distillers grains (particularly wet distillers grains with solubles) have all been associated with *E. coli* O157 prevalence in cattle feces. Results, however, are often conflicting, and other factors likely play a role. The factors that ultimately led to the emergence of STEC as a new pathogen in the 1980s have so far remained unclear.

**Antimicrobial-resistant non-STECE. coli**

**Reasons for inclusion: #1 and 2**

Antimicrobial-resistant *E. coli* other than STECs (“non-STECE. coli”) pose a distinct food safety issue, but their appearance in foods of animal origin is also an emerging public health concern.

Nonpathogenic *E. coli* are a natural and constant component of gut microflora and have a demonstrated ability to acquire, carry, and transfer resistance genes to pathogens that may be present in animals, the environment, or the human intestinal tract. The presence of drug-resistant nonpathogenic *E. coli* on meat or poultry products may therefore conceivably lead to the transmission of resistance genes to other bacteria, including potential pathogens, present on these foods or, after ingestion, inside the human gut.

Notably, even though STECs are among the pathogenic *E. coli* of greatest public health concern, various other *E. coli* strains are also associated with severe disease in humans, including extraintestinal pathogenic *E. coli* (ExPEC). For example, uropathogenic *E. coli* cause urinary tract infections (UTIs), and meningitis-associated *E. coli* are associated with meningitis and sepsis. *E. coli* strains can quite readily exchange genetic material, including antimicrobial resistance or virulence genes, and it is believed that the emergence of drug-resistant ExPEC infections, including recurrent UTIs, may be caused by foodborne exposure, particularly to retail poultry.
Emerging Antimicrobial Resistance

Assessing the relationship between antimicrobial use on farms and non-foodborne drug-resistant infections in humans is beyond the scope of this report, as well as a complicated issue: The horizontal transfer of resistance genes among unrelated bacteria; the spread of clonal strains among humans, animals, and the environment; and antimicrobial selective pressure from both human and animal use, make for a complex matter that cannot be easily dissected and studied.

Nevertheless, we know that agricultural antimicrobial use is associated with antimicrobial-resistant (AMR) bacteria in food animals, which can be shed into the farming environment and spread downwind or downstream. As a result, AMR bacteria can colonize or infect farmworkers, or those living nearby. As described elsewhere in the report, AMR pathogens can and do enter the food supply and cause human illness. The CDC estimates that drug-resistant Campylobacter and Salmonella, both primarily transmitted via foods of animal origin, are responsible for a combined 410,000 illnesses (i.e., 310,000 due to drug-resistant Campylobacter and 100,000 due to drug-resistant non-typhoidal Salmonella) and nearly 70 deaths (i.e., 40 due to drug-resistant non-typhoidal Salmonella and 28 due to drug-resistant Campylobacter) annually.

In the 1980s and 1990s, multidrug-resistant strains of Salmonella emerged, particularly among serotypes Typhimurium and Newport. In particular, multidrug-resistant Salmonella Typhimurium DT104 emerged as a global epidemic in both animals and humans. Since 1996, NARMS has been monitoring antimicrobial resistance among enteric bacteria isolated from humans, retail meats, and food animals. NARMS shows the proportion of human isolates of non-typhoidal Salmonella resistant to two or more classes of antimicrobial drugs has dropped since the late 1990s, although about 1 in 10 infections were still multidrug-resistant in 2013.††

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Disease burden

Analyses of *E. coli* isolates from humans and animals suggest food and animal sources for the colonization and infection of humans with antimicrobial-resistant *E. coli*. The presence of antimicrobial-resistant *E. coli* in food-producing animals, their environments, and their end products therefore likely has direct relevance for the human disease burden.

Many studies have documented antimicrobial-resistant *E. coli* in retail meats and poultry, sometimes at high prevalence. For instance, from 2002 to 2008, NARMS recovered and examined *E. coli* isolates for antimicrobial susceptibility from nearly 12,000 food samples representing retail chicken parts, ground turkey, ground beef, and pork chops. As shown in Table 2, NARMS found in 2013 that 75 percent of retail chicken samples, and 78 percent of ground turkey samples, were positive for *E. coli*. Moreover, they found that resistance profiles differed according to animal origin. More than half of turkey isolates (59 percent) were multidrug-resistant (defined as resistance to at least three drug classes), compared with 31 percent of chicken, 14 percent of pork, and 8 percent of beef isolates.
Table 2

Prevalence of Antimicrobial Resistance Among E. coli From Different Meats

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<thead>
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<tr>
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<tr>
<td>Overall E. coli prevalence</td>
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<tr>
<td>MDR &gt;= 5 drugs</td>
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<tr>
<td>MDR &gt;= 3 drugs</td>
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<td>2.5</td>
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<tr>
<td>Tetracycline</td>
<td>43.3</td>
</tr>
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</table>

* Based on data in Table 85a in the NARMS summary tables; Table 86a provides contradictory numbers.


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Emerging strains

Trends suggest that strains of *E. coli* present in meat and poultry are becoming more resistant to frontline drugs used for treatment in humans, including third-generation cephalosporins, fluoroquinolones, and trimethoprim-sulfamethoxazole.\(^\text{105}\)

Of particular public health concern among antimicrobial-resistant *E. coli* are strains that produce ESBLs, or extended-spectrum \(\beta\)-lactamases.\(^\text{106}\) These are a family of enzymes that confer resistance to third-generation (extended-spectrum) cephalosporins, as well as almost all other \(\beta\)-lactam antibiotics. ESBL genes in *E. coli* are typically found on mobile genetic elements, facilitating their carriage, accumulation, and transfer to other bacteria.\(^\text{107}\) ESBLs can be produced by a number of *Enterobacteriaceae*, but the most common are *Klebsiella* and *E. coli*. ESBL-producing *Enterobacteriaceae* are typically susceptible only to carbapenems and cephamycins, drugs of last resort. Carbapenem-resistant *Enterobacteriaceae* present one of the most pressing antimicrobial-resistance challenges facing human medicine today. The CDC estimates that health care-associated infections from carbapenem-resistant *Klebsiella* and *E. coli* are responsible for 9,000 infections and 600 deaths in the U.S. annually.\(^\text{108}\)

ESBL-producing *E. coli* have been documented in animals and retail meat, although the proportion of overall drug-resistant infections transmitted to humans by food is unknown.\(^\text{109}\) A Dutch study of ESBL-producing *E. coli* isolates found strong similarities in ESBL genes and multi-locus sequence typing (MLST) strains between isolates from hospitalized patients and retail chicken.\(^\text{110}\) This study is not likely representative of the U.S. but is indicative of the possibility of ESBL genes disseminating from animals to humans via food.

Changing exposure pathways

Epidemiological and microbiological evidence suggests that some exposures to ESBL-producing *E. coli* in food may have led to extraintestinal infections such as urinary tract infections. Of the 6 million to 8 million UTIs diagnosed in the U.S. each year, more than 85 percent are caused by ExPEC infections.\(^\text{111}\) In recent years, drug-resistant ExPEC have rapidly emerged, including those producing ESBLs, making treatment more difficult and less likely to succeed.\(^\text{112}\) In a number of unusual community outbreaks suggesting a common exposure source, foodborne exposure was suspected but never confirmed.\(^\text{113}\)

The potential role of food as a reservoir or source for these bacteria is supported by a number of studies finding ExPEC on chicken, turkey, pork, beef, and other foods.\(^\text{114}\) Some of these studies have identified close genetic similarities between strains from food animals, particularly chicken, and clinical isolates.\(^\text{115}\) A case-control study of women with UTIs found that higher rates of chicken consumption were associated with multidrug-resistant infections than with infections with strains that are susceptible to all antimicrobials.\(^\text{116}\) Furthermore, strong genetic similarities and similar antimicrobial resistance patterns have been documented between ExPECs and avian pathogenic *E. coli* (APEC), which causes extraintestinal colibacillosis in chickens.\(^\text{117}\)

ESBL genes have been detected in APEC and the fecal *E. coli* of healthy poultry.\(^\text{118}\) Some studies suggest that the drug-resistance of APEC and poultry-associated ExPEC strains may be a result of antimicrobial use, for example cephalosporins, that may be administered to prevent or treat poultry diseases.\(^\text{119}\) There are too little data, however, to systematically assess risk factors for ESBL emergence and evaluate potential associations with antimicrobial use on farms.

As with cephalosporin-resistant *Salmonella*, the most promising starting point for controlling ESBL-producing *E. coli* in poultry and other food animals may be to limit the use of cephalosporins, as was recently proposed by the
EFSA Panel on Biological Hazards, and as implemented by the FDA. Still, because ESBL-producing *E. coli* strains have been found in the environment and isolated from wildlife, stopping cephalosporin use on farms may not be sufficient to reduce carriage in food animals.

**Listeria monocytogenes**

**Reasons for inclusion: #1 and 2**

Although *L. monocytogenes* was first isolated from animals in the 1920s, its relevance to human health did not begin to be understood until 1949, as the result of an epidemic among newborns in Germany. However, questions about its epidemiology and transmission routes remained. During the 1980s, after a number of outbreaks in Canada and the United States, *L. monocytogenes* became recognized as a prominent foodborne pathogen. The risk of infection linked to the consumption of certain foods such as ready-to-eat meat appears to be declining, but an aging population will likely mean an increased public health impact in the future.

**Disease burden**

Foodborne listeriosis is a rare but serious infection, causing about 1,600 illnesses and 250 deaths annually in the U.S. Infections with *Listeria monocytogenes* are the third-leading cause of foodborne deaths in the U.S., behind *Salmonella* and *T. gondii*. Listeriosis has the highest costs of illness per case among all major foodborne pathogens: The estimated annual burden equals $2.8 billion a year, and the average cost per case has been estimated at $1.8 million.

The overall incidence rate of listeriosis was estimated to be roughly 0.8 cases per 100,000 people in 1989, and more recent FoodNet data from 2004 to 2009 suggest a rate between 0.25 and 0.32 cases per 100,000. The number of recorded outbreaks has increased, however, from about one every four years prior to the creation of PulseNet in 1998 to about 2.2 outbreaks per year since then.

Listeriosis is almost exclusively a disease of high-risk population subgroups, including the elderly, pregnant women and their newborns, and people with immune-compromising conditions. Within these high-risk population subgroups, listeriosis can cause a variety of severe clinical symptoms including septicemia, meningitis, and encephalitis. During pregnancy, listeriosis can result in miscarriage, stillbirth, preterm birth, septic conditions in newborns, and severe acute diseases with a heightened risk of serious neurodevelopmental sequelae. About 20 percent of symptomatic listeriosis cases result in death.

**Evolving understanding of strain-specific differences in public health risk**

Although all strains of the species *L. monocytogenes* are considered pathogenic, experimental and surveillance data suggest that certain subtypes pose a greater threat to public health than others. Of the 13 currently known *L. monocytogenes* serotypes, only three (1/2a, 1/2b, 4b) are responsible for over 95 percent of human infections; of these, serotype 4b has been mainly associated with large outbreaks, while serotype 1/2a has been mostly associated with sporadic cases. Serotype 4b has a higher reported hospitalization and case-fatality rate than 1/2a or 1/2b.

**Changing exposure pathways**

*L. monocytogenes* is a ubiquitous environmental bacterium, part of the fecal flora of many mammals, and widely distributed in many environments. It is also hardy, able to survive and grow in many conditions, including a wide range of pH values, high salt concentrations, and cold temperatures. Therefore, *L. monocytogenes* can
persist in cool, wet environments such as those experienced in some food manufacturing plants as well as most refrigerators.

The foods responsible for foodborne listeriosis have changed over the past 30 years, although some risks remain the same. A large Mexican-style cheese outbreak occurred in California in 1985. Over 140 people were infected, including 93 pregnant women, and 48 deaths occurred: 30 fetuses or newborns and 18 adults. Today, soft unripened cheeses remain a leading cause of listeriosis outbreaks.

Contaminated ready-to-eat meats have been another major source of listeriosis outbreaks in the U.S., although these appear to have peaked in the early 2000s and may now be declining. The decline is likely a direct result of concerted efforts by the food industry and regulatory agencies. Major successes have been achieved over the past decade in reducing contamination of deli meats through improved manufacturing practices in processing plants that minimize the risk of cross-contamination and in increasing formulation of products with substances that limit or prevent \textit{L. monocytogenes} growth. USDA-FSIS and others have found the risks from retail-sliced deli meats to be up to five times greater than those from prepackaged cold cuts. In addition, simulation studies have shown that limiting environmental cross-contamination and preventing \textit{L. monocytogenes} growth on contaminated products (e.g., through formulation with growth inhibitors or strict refrigeration at adequate temperatures) are the most promising interventions for reducing the risk of listeriosis associated with products handled at retail.

Changing population susceptibility

Exposure to \textit{Listeria monocytogenes} is believed to be quite common, however, most normal, healthy adults will not show clinical symptoms. Members of at-risk population subgroups, though, such as the elderly, pregnant women, and immunocompromised individuals, can suffer severe, potentially fatal illnesses. Based on U.S. surveillance data from 2004 to 2009, the incidence of listeriosis in the U.S. in the total population equals about 0.27 cases per 100,000 people, compared with 3.42 cases per 100,000 pregnant women and 1.21 cases per 100,000 individuals older than 65.

Although the incidence of listeriosis is not increasing, emergence remains a future concern. Because listeriosis risk increases with age and the U.S. population is aging (i.e., the proportion of elderly individuals is increasing), there are concerns that the incidence of listeriosis could increase in the short to medium term. The number of people 65 and older is expected to increase by 135 percent between 2000 and 2050, with the subpopulation of those 85 and older expected to increase by almost 350 percent.

Non-typhoidal \textit{Salmonella}

\textbf{Reason for inclusion: #2}

While \textit{Salmonella} is not a new pathogen, new strains are emerging—including some that are resistant to antimicrobial drugs. In addition, our understanding of food vehicles associated with foodborne \textit{Salmonella} infection is changing.

Disease burden

With very few exceptions (e.g., serotypes \textit{gallinarum} and \textit{pullorum}), non-typhoidal \textit{Salmonella} serotypes are zoonotic (i.e., capable of infecting animals and humans), even though individual serotypes differ in host range and some (e.g., serotype Dublin) are primarily associated with animals. Non-typhoidal \textit{Salmonella} strains are responsible for acute gastroenteritis, are primarily foodborne, and have been around a long time as well.
Recognition of their importance to human public health grew in the 1950s, and the CDC has been tracking *Salmonella* infections since 1962.\textsuperscript{142}

*Salmonella* is the leading cause of foodborne bacterial disease in the United States today, and the CDC estimates that this pathogen alone is responsible for over a million illnesses, 20,000 hospitalizations and nearly 400 deaths a year.\textsuperscript{143} The approximate annual cost of illness exceeds $3.7 billion.\textsuperscript{144}

Progress in reducing salmonellosis has been inadequate. In 1997, the annual reported incidence rate for *Salmonella* was 13.6 infections per 100,000 people. That rate became the baseline for the federal government’s Healthy People 2010 program, which set a target infection rate of 6.8 for 2010. By 2010, however, the reported incidence rate from FoodNet exceeded 15.\textsuperscript{145} Healthy People 2020 set a revised target rate of 11.4 cases per 100,000. However, the reported rate for 2013 was 15.9 infections per 100,000 people per year,\textsuperscript{146} about 17 percent higher than in 1997 and far above the new target rate.

**Emerging strains**

**Changing serotypes**

The somewhat static picture of overall salmonellosis incidence belies a dynamic picture of the strains responsible for disease. Hundreds of *Salmonella* serotypes are associated with human illness, many of which occur quite infrequently. Moreover, while some strains are in decline, others are emerging as important public health threats.

Figures 10 and 11 show incidence trends for the most common *Salmonella* serotypes. Figure 10 shows trends in the most commonly reported serotypes isolated by CDC’s FoodNet in 2014. Data are provided from 1999 to 2014 and are relative to the 1996-98 reference period. Figure 11 shows the relative change in the top *Salmonella* serotypes isolated in 2014, relative to 1996-98.

Both figures show declines in the frequency of *S. Typhimurium* and *S. Heidelberg*. Although *S. Typhimurium* had been the dominant serotype since surveillance began in the 1960s, its incidence has steadily declined; since 2007, *S. Enteritidis* has been the leading *Salmonella* serotype reported in lab-based surveillance and notifiable disease surveillance, and identified in outbreaks. Isolations of *S. Newport*, *S. Saintpaul*, and *S. Infantis* appear to be increasing. Likewise, in Figure 11, the best estimates for the incidence rates of *S. Muenchen* and *S. Oranienberg* show increases, though the confidence intervals for many of these are quite wide and cross the “no change” line, indicating that changes cannot be statistically proven.
Less common *Salmonella* strains are responsible for an increasing portion of reported human salmonellosis cases. Today, the leading serotypes reflect a lower proportion of overall isolates than they did in the past; the top five serotypes were responsible for about 65 percent of isolates submitted to CDC in 1987, but reflect only about 55 percent of isolates captured in the CDC’s Laboratory-Based Enteric Disease Surveillance (LEDS) system in 2011.\(^{147}\)

*Salmonella* serotypes differ, sometimes substantially, in pathogenicity and host range. *Salmonella* serotypes have been demonstrated to have different relative rates of invasive infection, and differ in the probability of severe outcomes, including hospitalization and death.\(^{148}\) Some emerging serotypes, for example Newport and Javiana, have relatively milder average courses of illness compared with leading serotypes like Typhimurium and Enteritidis, while others such as serotype 4,[5],12:i:- may be associated with worse clinical outcomes.\(^{149}\) There may also be variation in associated disease severity within serotypes;\(^{150}\) genetic methods of identification can be used to further differentiate strains within them.
Antimicrobial-resistant strains

Outbreaks of drug-resistant Salmonella are an emerging concern. Salmonella was the most common cause of antimicrobial-resistant foodborne outbreaks between 1973 and 2011 (48 of 55 outbreaks).151 Ground beef (n=10), poultry (n=7), and pork (n=1) together accounted for a considerable share of the 55 outbreaks with antibiotic-resistant bacteria.

Since 2012, there have been three additional notable foodborne outbreaks involving antimicrobial-resistant bacteria, all due to S. Heidelberg in chicken. One of the outbreaks sickened over 600 people and hospitalized more than 200 between March 2013 and July 2014. Of the 68 isolates tested during that outbreak, 44 (65 percent) were antimicrobial-resistant, of which more than half (i.e., 24 isolates) were multidrug-resistant.152

Drug-resistant Salmonella emerged as an international problem in the 1980s and 1990s. In particular, multidrug-resistant Salmonella Typhimurium DT104 emerged as a global epidemic in both animals and humans.153 Although the incidence of DT104 has been declining, other resistant strains have emerged.

Figure 11

Percentage Change in Incidence* of Culture-Confirmed Infections With the Top Salmonella Serotypes in 2014† Compared With Average Annual Incidence During 1996–98

* The estimates and confidence intervals presented indicate the relative change in the incidence of that serotype compared with 1996–98. The actual incidences of these infections cannot be determined from the graph.

† Data are preliminary.


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For example, the rise of cephalosporin-resistant *Salmonella* among both human and animal isolates is an emerging concern. Similarly, resistance to ceftriaxone (a related drug used as a first-line therapy to treat human salmonellosis and other infections, including infections with *Klebsiella* and *E. coli*).\textsuperscript{154}

Resistance to ceftriaxone, a related antibiotic, has also increased in human isolates over the same period.\textsuperscript{157} Across all serotypes of non-typhoidal *Salmonella*, resistance rose from 0.2 percent in 1996 to 2.9 percent in 2010, while increases were more pronounced for serotypes Typhimurium (from 0 percent to 4.7 percent), and Heidelberg (from 1.4 percent to 24.2 percent).

**Regulatory Response to the Emergence of Cephalosporin Resistance**

In response to concerns about the future efficacy of human cephalosporin therapy, the FDA first issued a broad ban on so-called extra-label\textsuperscript{*} uses of cephalosporin in 2008.\textsuperscript{†} This ban was revoked a short time later under criticism from the pharmaceutical and animal industries. In 2012, the agency issued a narrower ban on cephalosporins in animals, prohibiting most extra-label uses, including non-therapeutic uses, in cattle, swine, chickens, and turkeys.\textsuperscript{‡} The new rule does not cover the use of cephalaprin, an older cephalosporin drug not considered by the agency to contribute substantially to antimicrobial resistance development, or use to treat minor food-producing species such as ducks or rabbits. The rule also allows for limited extra-label use of cephalosporins in cattle, swine, chickens, and turkeys if it follows label instructions regarding dose, frequency of administration, duration of treatment, and administration route.\textsuperscript{§}

\* As specified in the Animal Medicinal Drug Use Clarification Act of 1994 (AMDUCA), veterinarians are, under certain circumstances, permitted to prescribe certain approved veterinary or human drugs for extra-label use, meaning in a manner not in accordance with the approved labeling. Extra-label use may deviate from labeled use in a variety of ways, as in the species to which the drug is administered, the disease or condition the drug is used to treat, or the dosage levels, frequency, or routes of administration.


\(‡\) Ibid.

\(§\) Ibid.
Canadian experiences with ceftiofur resistance support the view that ceftiofur use in animals may result in extended-spectrum cephalosporin resistance among bacteria isolated from animals and people. In Quebec in 2004, 62 percent of *Salmonella* Heidelberg isolates from chicken were ceftiofur-resistant, as were 36 percent of human isolates. In 2005, Quebec hatcheries voluntarily stopped the extra-label use of ceftiofur in eggs. By 2008, ceftiofur resistance among *S. Heidelberg* isolates dropped by two-thirds in both chickens and humans, to 18 percent and 12 percent, respectively. Ceftiofur-resistant *E. coli* in chicken dropped as well, from 34 percent in 2004 to 18 percent in 2008. Notably, corresponding reductions in the prevalence of resistance were also observed in neighboring jurisdictions. Whether this reflects indirect consequences of the ban in Quebec (e.g., shipment of eggs from Quebecois hatchers to neighboring provinces) or the impact of some external factors that affected multiple provinces has remained unclear.

**Changing exposure pathways**

Just as serotypes differ in associated disease severity, they also differ in transmission pathways and host range. Some *Salmonella* serotypes and subtypes are found solely in specific food animal species and their meat and animal products, while others are found in a wide variety of foods. Thus, serotypes have variable pathways to human exposure.

*S. Dublin*, for example, is almost entirely associated with cattle. Although this serotype is not a major cause of foodborne disease in the U.S., it has caused sporadic illnesses and outbreaks, and may be more likely to lead to severe clinical disease than other serotypes. Similarly, *S. Enteritidis* is primarily, yet not exclusively, associated with eggs and poultry, as are *S. Heidelberg* and *S. Hadar*.

Overall, poultry and eggs are considered the leading causes of foodborne salmonellosis based on case-control studies, outbreak analysis, and other epidemiological studies. A recent CDC study of foodborne outbreaks found distinct food attribution patterns for serotypes Enteritidis, Heidelberg, Newport, Javiana, Typhi, and Typhimurium, as well as for all others combined. While poultry and eggs cause the greatest share of illnesses caused by serotypes Typhimurium, Enteritidis, Heidelberg, and Hadar, the serotypes Newport and Javiana are largely associated with produce.

The findings from these and other studies point to an increasing role of food commodities other than meat, poultry, and eggs in foodborne salmonellosis. *Salmonella* can survive for long periods in secondary habitats outside of animal hosts, particularly aquatic environments. For example, *Salmonella* has been found to persist year-round and at high rates in surface waters of the southeastern United States. It also has been found to persist within agricultural environments despite efforts at decontamination. Moreover, flies and birds have been shown to be important vectors for transmitting *Salmonella* from farms to wildlife and local populations. More research, however, is needed to truly understand the complex natural cycles and interdependencies underlying environmental and foodborne exposure to *Salmonella*.

**Toxoplasma gondii**

**Reason for inclusion: #2**

*Toxoplasma gondii* is a parasite that was first identified in 1908, although its complete life cycle was not understood until 1970. In recent years, toxoplasmosis has been of emerging concern because foodborne transmission is increasingly recognized as a primary exposure pathway and because of rising concerns about public health impacts associated with latent infections.
Disease burden

*T. gondii*, a protozoan parasite, is one of the most universal human parasites on the planet, found in every human population that has been investigated. About one in five Americans have been found to have serological evidence of prior infection, although this rate is estimated to be in decline.170

The CDC estimates that foodborne toxoplasmosis kills over 300 people in the U.S. each year, making it the second deadliest foodborne pathogen in the United States behind *Salmonella*. Largely due to high rates of mortality among the most susceptible population subgroup (i.e., embryos and fetuses), the annual costs of illness associated with foodborne *Toxoplasma* is estimated at $3.3 billion, second only to *Salmonella*.171

In most people, infection is asymptomatic or mild, resulting in swollen lymph nodes, fever, malaise, aches and pains, and other flu-like symptoms. In the immunocompromised, however, disease manifestation is usually very severe, including encephalitis, myocarditis, and pneumonia. In these cases, death is almost certain without treatment. It should be noted that most such severe cases in immunocompromised individuals are believed to be due to reactivated latent infections, not new ones.

If a woman becomes infected during pregnancy, tachyzoites can be transmitted to the fetus, resulting in congenital infection (i.e., the baby is born infected). The likelihood of transmission is higher at later stages of pregnancy, but earlier transmission is associated with more severe disease. Estimates of the rate of congenital toxoplasmosis vary widely, from 1 to 100 cases per 100,000 births, equalling up to 4,000 cases per year.172 Many of these children are asymptomatic at birth, although they can develop chorioretinitis later in life. Miscarriage or neonatal death can also occur, as can severe outcomes such as blindness, mental retardation, and other central nervous system abnormalities.

To correctly understand infection risks it is important to understand the complex life cycle of this pathogen. *T. gondii* has three primary life-cycle stages: oocysts, tachyzoites, and bradyzoites. (See Figure 12.) Sexual reproduction occurs solely in cats, where oocysts are produced and shed with the feces in very high numbers, during brief periods of time. After one to five days, these oocysts become infective and can survive for 12 to 18 months in the environment. Sporulated (infective) oocysts ingested by a mammal (e.g., rodent, pig, or sheep) or bird transform into tachyzoites, a rapidly dividing stage that spreads infection inside this intermediate host. Eventually, tachyzoites localize in skeletal muscle, heart, brain, and eyes, where they convert to tissue cysts, or bradyzoites. The parasite remains in these tissues indefinitely, suspended in a largely inactive state. If the bradyzoites are ingested by a cat, infection is followed by sexual reproduction and the cycle begins anew. However, when tissue cysts in meat are ingested by “accidental hosts” such as humans, bradyzoites transform back into tachyzoites, infecting the new host, but sexual reproduction cannot follow. The life cycle is thus interrupted.

Humans can become infected through direct contact with cat feces or fecally contaminated food (e.g., fruits, vegetables); they can also become infected with bradyzoites by ingesting meat or offal from infected animals, such as undercooked pork or lamb. In addition, humans can be infected through blood transfusions or organ transplants from infected individuals, or from a pregnant woman to the fetus.

Emerging health impact

In the past decade, new and controversial research suggests that latent *T. gondii* infection in the brain may affect human personality and behavior. For example, a number of studies have associated schizophrenia with prior infection by *T. gondii*.173 The illness has also been associated with higher rates of depression, suicide, risky
Figure 12
Life Cycle of *Toxoplasma gondii*

Cat ingests tissue cysts and sheds oocysts

Herbivors ingest oocysts

Tissue cysts in the muscle of infected animals

Fecal oocysts

Ingestion of contaminated fruits and vegetables

Raw or undercooked meat

Contact with cat feces

Human infection

behavior, personality shifts, changes to physical appearance associated with testosterone and estrogen levels, cognitive deficits, and reduced concentration.174

Much more research is needed to verify and solidify these associations and establish causality. If \textit{T. gondii} infection does in fact contribute to any or all of these outcomes, the disease burden associated with foodborne toxoplasmosis, which is already quite high, could substantially increase.

Changing exposure pathways

The historic picture of \textit{T. gondii} infection was that of direct contact with cat feces, typically through a litter box or soil. However, over the past 15 years scientists have begun to understand the critical role of foodborne exposure. The CDC now estimates foodborne exposure to be responsible for about half of all \textit{Toxoplasma} infections in the United States each year.175

Foodborne exposure in people occurs from the ingestion of tissue cysts (bradyzoites) in raw or undercooked meat, tachyzoites in unpasteurized milk, or sporulated oocysts on plants or other foods that have been contaminated via water or soil. Routes of exposure are difficult to pin down in part because most acute infections are mild or asymptomatic.

Primary vehicles for foodborne transmission are estimated to be the ingestion of tissue cysts from raw or undercooked pork, lamb, and wild game.176 Unpasteurized goat’s milk, raw ground beef, raw shellfish, and fecal-contaminated soil on raw, unwashed fruits and vegetables have also been identified as risk factors.177 Numerous epidemiological studies have found undercooked meat to be an important risk factor, including a study that uncovered toxoplasmosis rates twice as high among meat eaters as among vegetarians.178 Surveys of retail pork, poultry, and beef have found low rates of contamination, although the prevalence of \textit{T. gondii} has been found to be quite high in meat from other animals such as sheep, lamb, and organically raised pork and chicken.179 A recent study in Wisconsin found that children who live on farms are five times as likely to show signs of previous infections (i.e., antibodies) as those who do not live on farms.180

The importance of \textit{T. gondii} in meat and poultry is likely substantial, but it is hard to measure. The overall incidence of \textit{T. gondii} infection is uncertain, as are proportions due to consumption of meat, poultry, or the numerous other routes of exposure.

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The CDC now estimates foodborne exposure to be responsible for about half of all \textit{Toxoplasma} infections in the United States each year.

**EPs with potential transmission through meat and poultry**

This section discusses the remaining pathogens identified in Table 1 as previously emerged or as potential emerging hazards to the U.S. meat and poultry supply. These bacterial, viral, and parasitic pathogens, as well as atypical prions, were chosen for one of three reasons (see Table 1 and the corresponding methodology text box for a full rationale of inclusion criteria):

**Reason #1:** They have been recognized as foodborne pathogens only since 1970181 and may therefore contain valuable lessons learned.
Reason #2: Expert panels convened by the European Food Safety Authority identified them as emerging or re-emerging diseases.\(^{182}\) (A comparable expert panel convened by U.S. authorities was not available at the time the report was written.)

Reason #3: They were identified as potential emerging threats by the experts involved in drafting this report.

For some of these pathogens, the potential for transmission with meat and poultry consumption has remained unclear or controversial. Other pathogens are not present in the U.S. but may conceivably be introduced in the country at a future time. These important caveats are highlighted where applicable. Notably, this section includes definitive and potential foodborne pathogens as well as pathogens that may be transmitted through contact with meat and poultry, such as MRSA.

**Bacterial pathogens**

*Arcobacter butzleri*

**Reason for inclusion: #1**

*Arcobacter* is a relatively newly identified genus of bacteria, related to *Campylobacter*. *A. butzleri* (formerly referred to as *C. butzleri*) is the most prominent of three species of *Arcobacter* associated with human enteric disease. *A. butzleri* shares similar microbiological and clinical features with *Campylobacter jejuni* but is more often associated with persistent, watery diarrhea than bloody diarrhea.\(^{183}\)

General foodborne infection has almost never been proven for *A. butzleri*, but it has been isolated from meats including raw chicken, pork, beef, and lamb, with the highest prevalence among them in chicken.\(^{184}\) *A. butzleri* was only recently identified as the likely cause of a 2008 gastroenteritis outbreak after consumption of chicken at a wedding reception in Wisconsin.\(^{185}\) Notably, targeted tests for *Arcobacter* species were explored only after the comprehensive culture and polymerase chain reaction (PCR) testing of five stool samples failed to identify more common pathogens, emphasizing the potential for underdiagnoses of *Arcobacter* and other emerging or uncommon pathogens.

Like many of the emerging *Campylobacter* species, pathogenic *Arcobacter* may play an important role in many outbreaks and cases of sporadic disease for which an etiologic agent is not identified.

*Clostridium difficile*

**Reason for inclusion: #3**

*Clostridium difficile* is an emerging gastrointestinal pathogen. Although foodborne transmission is not confirmed, meat and poultry may serve as an important pathway for infection, as discussed below. *C. difficile* has been isolated from food animals and from retail meats and poultry, among other foods.\(^{186}\)

*C. difficile* has historically been associated with hospital-acquired gastrointestinal infection, of which it is now the leading cause.\(^{187}\) Symptomatic *C. difficile* infections (CDI) generally follow disruptions to the intestinal microbiome. These interruptions are most commonly caused by treatments with antimicrobials, but they can also be the result of medical procedures, antacids, and more. Pathogenic strains of *C. difficile* can then overrun the gut and release toxins that cause bloating, diarrhea, and severe abdominal pain, sometimes with severe inflammation of the colon. These symptoms may last for weeks or months.\(^{188}\) Although treatment with antimicrobials can be successful in limiting symptoms, as many as 25 percent of patients have reported recurrences.\(^{189}\)
*Clostridium difficile* is a prototypical example of an EP: Disease incidence was stable until the mid-to-late 1990s, but starting in 2000 many studies have documented increasing CDI incidence, disease severity, recurrence, and antimicrobial resistance. At the same time, CDI is increasingly recognized as a cause of diarrhea not only in hospitals but also in communities, where it is affecting younger people and those without traditional CDI risk factors. More than 20 percent of CDI is now estimated to be community-associated, with some studies reporting a rate over 40 percent.

CDIs were once believed to be almost entirely spread through person-to-person transmission via hands, shared hospital rooms, or other surfaces. More recent research, including examinations of hospital outbreak isolates using whole-genome sequencing and MLST, has found great diversity and nonclonal strains (i.e., not all strains being genetically essentially identical), thereby undermining this transmission model. Many infections previously thought to be nosocomial (i.e., hospital-acquired) may have actually been caused by foodborne exposure through hospital meals or via asymptomatic carriage of *C. difficile* into the institution from the community.

Estimates of the share of adults who are asymptomatic carriers of toxigenic *C. difficile* vary from less than 2 percent to more than 7 percent, with causes of infection largely unknown. Potential risk factors for community-acquired CDI include asymptomatic carriers, use of proton pump inhibitors and other acid suppression medication, animal contact, and foodborne and waterborne exposure.

Although colonization and infection via foodborne exposure has not been firmly established, *C. difficile* has been isolated from the feces of production animals, retail meats, poultry, seafood, and vegetables. Some, but not all, molecular studies have found similar strains in humans and animals, suggesting zoonotic transmission or a shared environmental reservoir. Among the most common strains found in food are PCR ribotype 078 and PCR ribotype 027, important causes of community-acquired CDI. Further research is needed to characterize the role meat and poultry consumption plays in CDI infections.

**Helicobacter**

**Reason for inclusion: #3**

Like *Arcobacter*, the genus *Helicobacter* was differentiated from *Campylobacter*, in this case in 1989. Over 35 *Helicobacter* species have been identified, with more likely to be discovered imminently. The most prominent species is *H. pylori*, for which humans are the reservoir. It is harbored in the upper gastrointestinal tract by as much as half of the world’s population and is a known cause of gastric ulcers and stomach cancer.

Research into the role of *H. pylori* and other *Helicobacter* species in clinical human disease has expanded in recent years. Species other than *H. pylori* have been found in the gastrointestinal tracts and hepatic systems of other animals, and some have been associated (to varying degrees) with animal or human disease or both. Foodborne exposure, while hypothesized, has so far not been proven. For example, *H. pullorum* has been isolated from broiler chickens and laying hens and has been isolated from human patients with and without gastroenteritis. Still, for the most part, evidence linking *pylori* and non-*pylori Helicobacter* infection to human disease is limited. Far more research is necessary to understand the potential risks posed by this bacterium.

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*Clostridium difficile* is an emerging gastrointestinal pathogen. Although foodborne transmission is not confirmed, meat and poultry may serve as an important pathway for infection.
Methicillin-resistant Staphylococcus aureus

Reason for inclusion: #3

Like C. difficile, invasive methicillin-resistant Staphylococcus aureus (MRSA) is a serious and emerging public health threat with potential but unconfirmed foodborne exposure. Transmission between food production animals and people has been substantiated, and MRSA has been isolated from a wide variety of meat and poultry products. Notably, the risk of MRSA infection is distinct from the risk of S. aureus food poisoning, a common food safety problem recognized since the 19th century. The latter occurs if certain S. aureus strains capable of generating toxins contaminate and multiply in food under the right conditions for toxin formation (e.g., temperature and food environment). The clinical symptoms of S. aureus food poisoning depend on the toxin and the ingested dose, but are usually relatively mild gastrointestinal symptoms that spontaneously resolve within 24 to 48 hours without requiring specific treatments.

By contrast, invasive MRSA can cause skin and wound infections, or more severe illnesses such as pneumonia and bloodstream infections resulting in sepsis. An estimated 80,000 MRSA infections occurred in the United States in 2011, resulting in over 11,000 fatalities.

In addition to hospital-associated strains, MRSA has found a reservoir in livestock, particularly swine, as well as horses, cattle, dogs, and cats. MRSA subtype 398 (ST398) was discovered in the early 2000s and has emerged as an important cause of human infection, often associated with exposure to livestock. A large study involving more than 30 authors from 20 institutions employed molecular methods on a diverse set of 89 ST398 isolates from humans and animals in 19 countries. The authors determined that this lineage originated as methicillin-sensitive S. aureus (MSSA) in humans, crossed the species barriers to pigs, and subsequently acquired tetracycline and methicillin resistance. Using state-of-the-art analyses, another group of researchers subsequently determined the existence of two distinct strains of MRSA ST398—one associated with humans and one with animals. The livestock-associated strain was found in a hospital environment and in newborn babies.

Numerous studies in the United States and elsewhere make a compelling case for transmission of MRSA strains, including ST398, from livestock to humans. Some of these studies have found higher colonization and/or infection rates of MRSA among farmworkers, veterinarians, and others in close proximity to animals, potentially indicating direct zoonotic transmission.

Of perhaps greater concern for public health are studies showing that proximity to animal production confers a risk of infection to the broader community. A recent study at an Iowa Veterans Affairs hospital found a threefold increased risk of MRSA colonization among people living within one mile of large swine facilities. Proximity to the application of swine manure has also been found to be a significant risk factor for MRSA infection.

MRSA is not typically considered a foodborne pathogen, although community-associated and livestock-associated strains have been isolated at high rates in recent years from retail meat in the U.S. and elsewhere. Even if food consumption does not develop as a significant risk factor, handling raw meat may pose a transmission risk for MRSA from the farm to the community.

Mycobacterium avium subspecies paratuberculosis

Reason for inclusion: #3

Mycobacterium avium subspecies paratuberculosis (MAP) is the etiologic cause of paratuberculosis in ruminants, a fatal gastrointestinal disease that largely affects cattle. It is also a suspected cause of Crohn’s disease in...
humans. Numerous epidemiologic studies have found associations between MAP and Crohn’s disease, but causality has not been confirmed and remains in dispute.

If MAP causes Crohn’s, it could have implications for the safety of meat, milk, and dairy products. MAP is widespread in dairy herds, where it is shed in milk and feces; it has been found in milk supplies and in high concentrations can survive commercial pasteurization and cheese production processes. It has also been detected on carcasses and in organ meats. Because fecal shedding of MAP is much higher in older animals, risks of meat contamination are higher in the carcasses of culled dairy cows (i.e., cows that have been removed from the herd and sent to slaughter) than from younger animals (less than 3 years old) raised for meat. Limited studies have found that some MAP cells may survive cooking.

**Yersinia**

**Reason for inclusion: #1**

The genus *Yersinia* spp. comprises pathogenic and nonpathogenic species. Among the human pathogens are enteropathogens such as *Y. enterocolitica* and *Y. pseudotuberculosis*. Infections of *Y. enterocolitica* and *pseudotuberculosis* primarily cause fever, diarrhea, and abdominal pain that can be misdiagnosed as appendicitis. Septicemia, skin rash, and other complications have been reported but are rare. Despite *Y. enterocolitica*’s first being isolated in 1934, its importance as a foodborne pathogen was not recognized until 1976. According to CDC estimates, approximately 90 percent of yersiniosis cases in the U.S. are foodborne. An estimated 97,656 foodborne yersiniosis cases are acquired in the U.S. each year. One study (based on expert elicitation) showed approximately 72 percent of U.S. yersiniosis cases attributable to pork, followed by 12 percent attributed to dairy and 5 percent to seafood.

*Y. enterocolitica* is not emerging in the U.S. Incidence of infection with *Y. enterocolitica* has declined from a rate of about 1 in 100,000 in 1996 to about 0.28 in 2014. It was included in this report because of the potential lessons learned. Its recognition as a foodborne pathogen as late as 1976 demonstrates the fact that even for important foodborne pathogens the role of foodborne transmission may remain unclear for years or even decades. *Y. enterocolitica* is believed to be a commensal organism (i.e., an organism that benefits from another organism without affecting it either positively or negatively) of swine, so as the feral swine population increases in density and expands in range, *Y. enterocolitica* could conceivably become a larger risk to public health in the future.

**Viruses**

**Avian influenza virus**

**Reason for inclusion: #2**

Avian influenza virus (AIV) is primarily a veterinary public health concern even though instances of foodborne transmission have been documented. A variety of influenza viruses can infect poultry as well as wild and pet birds. These viruses are typically classified as either low pathogenicity (LPAI) or high pathogenicity (HPAI) based on the disease they cause in poultry.

HPAI viruses result in large outbreaks and severe illness, with very high mortality rates in poultry flocks. In fact, between December 2014 and June 2015, the U.S. experienced a devastating HPAI outbreak that severely affected parts of the U.S. turkey and egg industry. LPAI viruses can also cause outbreaks among poultry but are generally not characterized by severe clinical disease. However, through spontaneous genetic changes, LPAI
viruses can develop high pathogenicity. Most HPAI and LPAI viruses appear to be largely apathogenic in wild birds, which are believed to be an important reservoir for poultry infections and may disseminate the virus along migration routes.\textsuperscript{236}

Al viruses generally do not infect humans; however, human-pathogenic Al viruses of the H5N1 and H7N9 subtypes have been emerging since 2003.\textsuperscript{237} These infections can be severe and potentially fatal. Human cases appear to be primarily sporadic and have been reported in Canada and several countries in Africa, southeast and central Asia, and the Middle East.\textsuperscript{238}

Many questions remain about the epidemiology, transmission dynamics, and foodborne risks associated with these human-pathogenic strains. In general, contact with live or dead birds, their respiratory secretions, blood, feces, organs, or their environment is believed to be the primary risk factor for human infection, with a few cases also linked to consuming food made from raw poultry blood.\textsuperscript{239}

Starting in December 2014, several HPAI strains have caused avian influenza outbreaks in commercial and backyard poultry flocks as well as wild bird populations in the U.S. While none of these strains are believed to be human-pathogenic, the outbreaks demonstrate the risk of HPAI virus introduction.\textsuperscript{240}

Crimean-Congo hemorrhagic fever virus

\textbf{Reason for inclusion: \#2}

Crimean-Congo hemorrhagic fever virus (CCHFV) is the cause of a potentially fatal hemorrhagic fever in humans, with a fatality rate of 10 to 40 percent.\textsuperscript{241} CCHFV occurs sporadically in Africa, Asia, the Middle East, and parts of Europe, and it may be emerging or re-emerging in several geographic regions including India and parts of southern Europe.\textsuperscript{242}

CCHFV is a tick-borne virus in the family \textit{Bunyaviridae}.\textsuperscript{243} A variety of domestic animal species, including cattle, sheep, and goats, are susceptible to infection but are not believed to exhibit clinical symptoms.\textsuperscript{244} Human infections are thought to occur primarily as a result of contact with the blood or tissues of infected livestock, or from the bite of an infected tick.\textsuperscript{245} A variety of tick species are believed to be vectors for CCHFV.\textsuperscript{246} Many epidemiological aspects associated with the establishment of disease foci, and in particular with a limited geographic spread in some areas, remain unclear.\textsuperscript{247}

The potential for foodborne transmission of CCHFV is currently unknown, but risk is generally believed to exist through contact with meat or organs, at least immediately after slaughter.\textsuperscript{248} There have been no known domestically acquired CCHFV infections in the United States, but the possibility for its introduction to the U.S. cannot be excluded.

Hepatitis E virus

\textbf{Reasons for inclusion: \#1 and 2}

Hepatitis E virus (HEV) is a cause of acute hepatitis in humans.\textsuperscript{249} While many infections are believed to be self-limiting, HEV infection can lead to liver failure, particularly in pregnant women and patients with chronic liver disease. It can cause chronic hepatitis, primarily among immunosuppressed individuals.\textsuperscript{250} In addition, infection during pregnancy can lead to premature birth, fetal loss, and health problems for the newborn such as hepatitis or hypoglycemia.\textsuperscript{251}

HEV is a small, non-enveloped virus with a positive-sense, single-stranded RNA genome.\textsuperscript{252} Four genotypes are
HEV appears to be endemic among swine in the U.S. and other countries, and it has been isolated from commercial pig livers sold in U.S. grocery stores. In 2007 and again in 2013, outbreaks of HEV in France were attributed to the consumption of raw pork liver sausages.

currently recognized—HEV-1, HEV-2, HEV-3, and HEV-4—all of which infect humans, and the distinction of additional genotypes has been proposed.253 The first two types, HEV-1 and HEV-2, generally cause more severe clinical symptoms.254

HEV was first isolated from swine in the U.S. in 1997, and infection is believed to be asymptomatic.255 Despite numerous questions about epidemiology and transmission dynamics, the zoonotic nature of HEV-3 and HEV-4 is generally accepted, and the pathogen has been isolated from a variety of animal species, including swine, cattle, sheep, deer, and rabbits.256

Potential foodborne transmission of HEV from pork and pork products has received increased attention in recent years.257 The virus appears to be endemic among swine in the U.S. and other countries, and it has been isolated from commercial pig livers sold in U.S. grocery stores.258 In 2007 and again in 2013, outbreaks of HEV in France were attributed to the consumption of raw pork liver sausages.259

Many questions remain about HEV’s potential foodborne transmission through pork or other meats. However, if a foodborne risk is firmly established, HEV could present an emerging global food safety risk.

**Middle East respiratory syndrome coronavirus and related coronaviruses**

**Reason for inclusion: #2**

Middle East respiratory syndrome coronavirus (MERS-CoV) causes a primarily respiratory disease in humans260 and is emerging in the Middle East. Infection can lead to pneumonia, respiratory failure, and in some cases other symptoms, including diarrhea.261 Current World Health Organization (WHO) estimates place the case-fatality rate up to 36 percent,262 though numbers may be skewed by the underdiagnosis of milder infections.

MERS-CoV is a small, enveloped virus with RNA genome; it is a relative of SARS-CoV, which caused the severe acute respiratory syndrome (SARS) pandemic of 2003.263 MERS-CoV was first isolated from individuals with acute respiratory illness in Saudi Arabia in 2012.264 As of June 2016, 1,768 laboratory-confirmed cases had been reported to the WHO, including at least 630 deaths.265 They were reported from various countries in the Middle East, and some presumably travel-related cases have been reported in the United States, France, Germany, Italy, the United Kingdom, Malaysia, the Philippines, and elsewhere.266

Many aspects concerning the emergence, epidemiology, transmission dynamics, and host range of MERS-CoV so far remain unclear. However,
serological and virological evidence suggests that the virus may have originated in bats, that dromedary camels may be an important source of human infections, and that the virus may have been circulating in the camel population for as long as several decades. Dromedary camels have been shown to shed MERS-CoV in their feces and milk, and people who work closely with camels appear to be at a heightened risk of infection. Whether the consumption of undercooked camel meat poses a transmission risk remains to be determined. Serological surveys of other animals, including goats, sheep, and cows, have as yet provided no evidence of infection.

MERS-CoV does not seem to pose a threat to the U.S. meat supply. However, should host range and geographic distribution change, MERS-CoV may pose an emerging hazard in the future. Notably, it has been theorized that the related SARS-CoV entered the human population through foodborne exposure, including consumption of civet cats, an intermediate host, and potentially bat meat.

**Rift Valley fever virus**

**Reason for inclusion: #2**

Rift Valley fever virus (RVFV) causes an illness in humans believed to manifest most often as a benign flu-like sickness. In rare cases, however, severe forms can be associated with meningoencephalitis, hemorrhagic fever, and retinal lesions, which may result in permanent loss of vision. RVFV, a member of the *Bunyaviridae*, is a small, enveloped virus with RNA genome. It can infect humans and a broad range of animals, including cattle, sheep, and goats. Mosquitoes and infectious wildlife are thought to be the primary sources of infection for livestock. For humans, consumption of meat and possibly raw milk have been proposed as potential transmission routes, with mosquito bites and contact with infected animals or their secretions another potential risk. RVFV is endemic in tropical regions of eastern and southern Africa, where it is believed to primarily circulate among ruminant wildlife reservoirs and hematophagous (i.e., blood-sucking) mosquitoes. Explosive outbreaks among livestock occur periodically (i.e., every five to 25 years), probably driven by a combination of climatic factors (i.e., precipitation resulting in explosive hatching of mosquito eggs) and demographic ones (i.e., availability of naive hosts that have not previously been in contact with the virus).

RVFV is not endemic in the U.S. However, a broad range of hematophagous mosquitoes, including certain *Aedes*, *Anopheles*, and *Culex* species, are believed to be capable of transmitting RVFV. The emergence of West Nile virus, a mosquito-borne *flavivirus* introduced in the U.S. in 1999 that has since become endemic, demonstrates the potential risk of introducing vector-borne diseases into the U.S. if a competent vector (i.e., one that can effectively transmit the disease) naturally occurs or is introduced.

**Parasites and prions**

**Cryptosporidium parvum**

**Reason for inclusion: #1**

Cryptosporidiosis has been recognized as a potential foodborne disease of humans and livestock since the 1970s and emerged as an important zoonotic human pathogen in the wake of the HIV/AIDS epidemic. *Cryptosporidium parvum* is a zoonotic parasite commonly shed by cattle and other livestock species. It causes diarrheal illness in humans as well as livestock, particularly neonatal calves and lambs, around the world. In immunocompetent individuals, this illness is typically self-limiting and non-life-threatening, but for those who are
immunocompromised, particularly HIV/AIDS patients, disease manifestations are severe and potentially fatal.\textsuperscript{285}
For these patients, severe extraintestinal manifestations such as biliary tract disease, respiratory infections, and pancreatitis are possible.\textsuperscript{286}

\textit{Cryptosporidium parvum} is believed to be transmitted primarily through the ingestion of food or water that has been contaminated with fecal material shed by infected livestock.\textsuperscript{287} Consumption of water and fresh produce presumably contaminated by livestock manure has been identified as a source of \textit{Cryptosporidium} exposure in various outbreaks and case-control studies, indicating a clear risk associated with manure runoffs.\textsuperscript{288}

Consumption of contaminated uncooked or undercooked meat may also pose some risk of infection. An expert elicitation\textsuperscript{289} study examining the sources of foodborne illness in the U.S. attributed 60 percent of cryptosporidiosis cases to produce, 9 percent to beverages, 7 percent to beef, and 6 percent to dairy.\textsuperscript{290}

**BSE and new variant Creutzfeld-Jacobs Disease (vCJD)**

**Reason for inclusion: #2**

Variant Creutzfeld-Jacobs Disease (vCJD), first described in 1996, is a rare, invariably fatal neurodegenerative disease that is presumably linked to the consumption of beef from cattle infected with bovine spongiform encephalopathy (BSE), even though the time between consumption and illness probably spans many years.\textsuperscript{291} A very small number of classic CJD cases used to occur prior to the emergence of vCJD, either as sporadic cases or linked to genetic predisposition or iatrogenic exposures (i.e., related to medical treatments).\textsuperscript{292} Multiple aspects distinguish vCJD from classical CJDs, including disease onset in younger patients (median death at 28 years compared with 68 years), a longer duration of disease, specific clinical symptoms, and distinctive neuropathological changes.\textsuperscript{293}

CJD and BSE are conditions classified as transmissible spongiform encephalopathies (TSEs); they have long been known to occur with low prevalence among humans (e.g., kuru among cannibals) and other species such as sheep and goats (i.e., scrapie), deer (i.e., chronic wasting disease), or mink (i.e., transmissible mink encephalopathy), and some risk of cross-species transmission may exist.\textsuperscript{294}

TSEs are generally believed to be caused by prions, “misfolded”\textsuperscript{295} proteins that do not contain genetic material but that can “replicate” by inducing other proteins to misfold, even though some scientific controversy remains about the causative agent, and many aspects related to pathogenicity, transmission dynamics, genetic predisposition, strain differences, and emergence are still unclear.\textsuperscript{296}

BSE was first diagnosed in U.K. cattle in the 1980s and presumably emerged through feed-borne exposure of cattle to infectious scrapies prions in meat and bone meal (MBM).\textsuperscript{297} BSE is a progressive neurodegenerative disease that primarily affects cattle and exhibits a long incubation period, averaging four to five years in cattle.\textsuperscript{298} Other livestock species, including sheep and goats, appear susceptible to infection, and intentional infections of laboratory animals by scientists indicate a potential for transmission to species such as pigs, cats, mink, and primates.\textsuperscript{299}

Feed-borne exposure to MBM is thought to be the primary transmission route for BSE among cattle, even though maternal transmission cannot be completely excluded.\textsuperscript{300} Control options implemented to contain the spread of BSE among cattle included a ban on the use of animal-derived proteins presumably at high risk for transmission (such as MBM) in bovine feed, the implementation of a passive animal disease surveillance system, and an active surveillance system based on the routine diagnostic testing of cattle at slaughter, paired with trace-back investigations on positive samples.\textsuperscript{301}
Diagnostic tests are more sensitive toward the end of the incubation period because by then considerable amounts of atypical prions (abnormal PrP) have been accumulated.\textsuperscript{302} Because of the long incubation period for BSE among cattle and the potential slaughtering of cattle in some production systems and geographic regions before diagnostic tests could detect BSE, incidence rates may not be comparable across countries or periods in time.\textsuperscript{303} The impact that changing concentrations of atypical prions during the incubation period may have on infectivity and the risk to human transmission is difficult to quantify.

The incidence of diagnosed BSE cases among livestock has been declining.\textsuperscript{304} Today, only a small number of cases are diagnosed in cattle in the U.K. and around the world, and the U.S. has been categorized by the World Organisation for Animal Health as having a negligible BSE risk.\textsuperscript{305}

Control options implemented to contain the spread of BSE among cattle included a ban on the use of animal-derived proteins such as meat and bone meal in bovine feed, the implementation of a passive animal disease surveillance system, and an active surveillance system based on the routine diagnostic testing of cattle at slaughter, paired with trace-back investigations on positive samples.

**Recommendations and conclusions**

**Emerging pathogen preparedness**

While today’s microbial risks in meat and poultry appear similar overall to what they were a decade or more ago, some specific and important differences exist. Food safety risks have not remained static; some microbial hazards, many parasitic diseases in particular, have been successfully controlled or eliminated, while some new microbial hazards have emerged. It is important to recognize that newly emerging hazards often are not well understood initially. The specific implications for food safety and public health may materialize almost immediately, as in the case of \textit{E. coli} O157:H7 in undercooked hamburgers, or may remain opaque for quite some time. The latter was, in fact, the case for some pathogens such as \textit{Listeria monocytogenes} that are now unanimously accepted as major food safety risks.

The emergence of new foodborne hazards can be driven by the microbe’s acquisition of virulence or resistance characteristics, by demographic changes and associated increases in population susceptibility, or by a pathogen’s emergence in a new host species or geographic region. In addition, new scientific knowledge may lead to re-evaluating and re-prioritizing of known risks.

Detecting the emergence of pathogens can pose tremendous challenges. Existing surveillance systems may not be adequate to detect EPs in a timely manner. Diagnostic tools may be incapable of distinguishing the EPs from any existing ones, and determining causal relationships between the presence of a pathogen and clinical disease can be extremely difficult. Sufficient epidemiological evidence may not be available to pinpoint transmission routes, especially if disease dynamics are complicated by multiple hosts and exposure pathways. The public health risks posed by the EP are often unclear, and data to assess these risks are scarce. Even if these challenges can be overcome, adequate regulatory policies are rarely in place to address the issue, and the relevant food industry segments may be ill-prepared to address the emerging hazards.
In this report, we discuss a number of emerging microbial hazards that may be transmissible through the U.S. meat and poultry supply. These include bacteria, viruses, parasites, and prions. For the purpose of emerging disease preparedness, these hazards can be broadly categorized into three distinct classes:

1. Completely new, previously unknown pathogens

The emergence of these zoonotic pathogens is an exceedingly rare event, but one with immense consequences for public health and the economy. This type of emergence is typically caused by a mechanism called “host jumping,” where a pathogen acquires the ability to infect a new species. For example, BSE emerged from scrapie or a related prion disease, and some avian influenza viruses acquired the ability to infect humans. Often, these emergence events are initially limited in geographic distribution, but they have the ability to expand quickly in geographic range. For example, the H5N1 avian influenza virus emerged in Asia and BSE emerged in the U.K., but both soon posed a pandemic threat.

Because these pathogens are truly “new” pathogens, many aspects related to epidemiology, transmission dynamics, and public health risk are initially unknown, and vaccines or treatments are not available. These emergence events tend to receive extensive public attention and media coverage. For instance, the emergence of SARS coronavirus in 2003 prompted the U.S. CDC to handle more than 10,000 press calls, 21 telebriefings, 23 health alerts, more than 2,000 clinical consultations, and three satellite broadcasts; there were eight laboratory-confirmed SARS cases in the U.S., all travel-associated, and no fatalities.

In many cases, when a new pathogen emerges, civic life is uprooted, trade is disrupted, and the economy is devastated. For instance, the emergence of Nipah virus, first recognized in Malaysia in 1998, caused at least 257 confirmed human infections and 115 fatalities, affected more than 1,000 pig farms, and led to the culling of more than 1 million pigs. Government compensations for the destroyed pigs alone equaled an estimated $35 million in U.S. dollars. The cost of the national control program equaled $136 million, and lost tax revenue has been estimated at $105 million. An estimated 36,000 people became unemployed. More than 700 homes and businesses were evacuated. Malaysian pork exports came to a halt, and local pork consumption dropped by up to 80 percent. Financial losses to the pork industry during the outbreak alone totaled an estimated $124 million.

Because of the considerable societal impact, these emergence events often lead to transformative regulatory changes. For instance, as a result of the Nipah outbreak, Malaysia now permits pig farming only in “identified pig farming areas.” After the emergence of human-pathogenic avian...
influenza viruses, at least some live bird markets in Indonesia implemented WHO recommendations to reduce the transmission risk. The emergence of BSE led to fundamental changes in cattle feeding and slaughter practices around the world, including restrictions on the repurposing of offal for ruminant feed and a prohibition on the slaughter of “downer cows” that are unable to walk.

Given the potentially enormous impacts these types of emergence events may have on all aspects of society, considerable research efforts have focused on trying to predict them. Certain risk factors, such as some pathogens’ broad host range, zoonotic nature, and “genetic plasticity” (e.g., high mutational rates or predisposition to horizontal gene transfer); predisposing climatic factors; health care system attributes; and structural factors (in particular those that affect contact rates between animals and humans) have been linked to an increased risk of emergence, but so far these types of emergence events have remained largely unpredictable.

Detecting EPs rapidly poses another challenge. For instance, evidence suggests that Nipah virus caused illnesses in humans and pigs for at least a year before it was recognized, and MERS-CoV may have been around in camels for decades before it was detected. However, technological advances such as next-generation (or high-throughput) sequencing and new phylogenetic methods, coupled with improved surveillance and collaboration among public and private entities around the world, allow EPs to be characterized and controlled more rapidly than ever. For instance, when a new, swine-origin, human-transmissible H1N1 influenza virus was isolated from two patients in California in March 2009, 45 public health laboratories in the U.S. were already using diagnostic methods capable of detecting the new strain. By late April 2009, swine were identified as the likely source of the outbreak strain, but evidence showed that transmission during the outbreak occurred person-to-person and that the initial outbreak occurred in Mexico. By April 21, CDC had begun developing a candidate virus for a new influenza vaccine, and by the end of the month, CDC was granted permission by FDA to manufacture and distribute diagnostic kits specific to the new strain. Within weeks kits began to be distributed to clinical and public health laboratories in all U.S. states and in more than 150 countries around the world. As new H1N1 viruses were isolated from case patients, information about the genetic makeup was made available to the public, starting in April 2009. Soon, independent academic researchers were able to confirm that the new virus indeed derived from several viruses that circulated in swine and that the initial transmission to humans had occurred months before the outbreak was recognized. On Sept. 15 of the same year, FDA announced the approval of several vaccines specific to the new H1N1 strain, and the first doses were administered in early October. Supplies were initially limited, but by the end of December, the vaccine was available to anyone who wanted it, although H1N1 activity had markedly decreased by then. Thus, even if it may not yet be possible to predict these types of emergence events, stopping them in their tracks may be a real possibility.

2. New strains of previously known pathogens

These strains typically emerge through the acquisition of virulence or antimicrobial resistance genes or both. In some cases, such emergence events were completely unanticipated and had a tremendous impact on public health, as in the case of the emergence of E. coli O157:H7 in ground beef. In many other instances, however, the emergence of new strains was perhaps more predictable. For instance, several new Salmonella strains with antimicrobial resistance have emerged over time, indicating a foreseeable pattern.

In many cases, the emerging traits have limited effects on the fundamental aspects of epidemiology and host range, and diagnostic methods are typically available to detect the pathogen, even though they may be inappropriate for discriminating an emerging strain from one that is more established. Regulatory policies and existing interventions, however, may be inappropriate for addressing the altered public health risk posed by the emerging strain, and the associated public health risks may not be fully understood.
While it may not be possible to predict the exact strain that will emerge, an understanding of the driving selection forces (e.g., antimicrobial use) and underlying genetic determinants may allow for a prediction of which pathogens and characteristics are more likely to emerge in the future. For example, various *Salmonella* strains with new antimicrobial resistance patterns have emerged in the past and are likely to continue to do so under current antimicrobial use practices.

3. Known strains to which susceptibility has been increasing, or that are expanding in geographic range

These emergence events are not directly driven by the pathogen or the food supply; driving forces may include increased immune suppression in the human population (e.g., due to aging, therapeutic drug use, or comorbidities) or an expanded geographic distribution of vectors, such as ticks for vector-borne diseases. The primary drivers here tend to be outside of the food chain. Existing diagnostic tools typically continue to be appropriate for the detection and characterization of the pathogen, and epidemiology and transmission dynamics are often relatively well understood. However, existing regulatory policies may be inappropriate for addressing the new risks. A thorough understanding of risk factors for infection and underlying population dynamics may allow the emergence of these pathogens to be more readily predicted than the preceding two types of EPs.

Regardless of the underlying mechanism, new microbial hazards will continue to emerge, and some of these will be transmissible through the meat and poultry supply. Improvements in basic research and risk assessment will allow prediction of certain emergence events, such as the emergence of new strains with new antimicrobial resistance or virulence patterns. Surveillance systems can be, and often already are, designed to detect these hazards as they emerge. Better surveillance across the food chain can expedite the detection of these hazards. Risk assessments, basic research, and related data such as on-farm antimicrobial use patterns can expedite and streamline the regulatory and industry response to emergence.

The emergence of completely new pathogens, however, is an extremely rare event with high impact but exceedingly low probability. Currently, it is not possible to predict emergence events ahead of time. However, general steps such as improvements in diagnostic capabilities and surveillance across the food chain will expedite the detection of these events, and risk assessments may prove useful in assessing the risk posed to consumers by a specific new pathogen. As recent history proves, rapid responses to such emergence events may be possible and may allow us to stop a new pathogen in its tracks.

New hazards will continue to emerge, and some of them will affect our food supply. Some we will be better able to prepare for than others. Improvements in surveillance, diagnostics, basic research, and risk assessment can help us be better prepared for the inevitable next emergence events. Being ready ahead of time will be crucial to allow for an expeditious, coordinated, and appropriate response to new emergence events.

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**Surveillance systems can be, and often already are, designed to detect new hazards as they emerge. Better surveillance across the food chain can expedite the detection of these hazards.**

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**Recommendations**

Preparing for disease emergence has to be premised on the fact that not every emergence event can be predicted or prevented, and not every emergence will be detected immediately. The desire to learn from the past has to be balanced against the understanding that every disease emergence event is different, shaped by conditions at a
certain place and time and by specific pathogens and animal species involved. The allocation of scarce resources to emerging disease preparedness has to acknowledge that some EPs go on to cause devastating pandemics that require global coordinated disease control efforts, while others remain restricted in geographic location and size. Yet predicting the magnitude of the associated public health problem is often extremely challenging. Therefore, the cost-effectiveness of any EP preparedness effort is nearly impossible to evaluate, as it depends on inherently unknown factors: the cost associated with the averted emergence event, the probability that the emergence in fact occurred, the cost of the implemented measure, and the potential opportunity cost of diverting scarce resources from other areas. More research to quantify the cost of previous emergence events as well as the cost of surveillance and the potential opportunity cost associated with the reallocation of scarce resources could be a first step toward evaluating the cost-effectiveness of EP preparedness. Nonetheless, as this report has shown, the economic, societal, and public health implications of an EP event can be devastating, as can the impact on the affected food industry as a result of eroded consumer trust. Therefore, EP preparedness is warranted.

As a first step toward preparedness for EPs, the U.S. should adopt a more holistic approach to food safety that directly acknowledges the dynamic and interconnected ecology of microorganisms, humans, food animals, and wildlife, and their respective environments. Such an approach will necessarily focus more heavily on the farm and feedlot than current ones do. Emerging disease preparedness should focus resources on multiple fronts:

1. Prediction
   a. Support efforts to understand what factors lead to the emergence of new pathogens, in order to identify clues for predicting and mitigating the risk of future disease emergence. Additional efforts should strive to analyze how regulatory and industry stakeholders have responded to such pathogens in the past to develop an understanding of what has, and has not, worked in different situations, and why.
   b. Support research (e.g., risk assessment, “systems thinking”) that, through conceptual approaches, helps to identify weaknesses in the current food production practices that may give rise to EPs.
   c. Identify and monitor trends in the food chain and beyond that may lead to disease emergence, such as demographic changes; changes in agricultural and animal husbandry; food manufacturing or processing changes; or changes in food preparation and consumption practices.

2. Detection
   a. Build surveillance systems and diagnostic tools that are able to detect EPs early, and that can reliably distinguish them from other microbes that occasionally occur but never pose a public health risk. This will allow for rapid response when necessary and minimize the risk of false alarms that waste resources and can erode public trust. Support efforts to determine when and where surveillance should be focused.
   b. Invest in collaboration with veterinarians, food industry scientists, and academics to improve detection, including syndromic surveillance and the use of innovative crowdsourcing approaches where applicable.

3. Capacity building
   a. Develop agile regulatory approaches, tools, and infrastructure (e.g., rapid risk assessments, expert panels) to foster quick responses in the face of large uncertainty. Develop approaches that will allow
for mitigation options to be implemented quickly and efficiently along the food chain, such as rapid approval processes for vaccines or therapeutics.

b. Invest in building the necessary relationships before a new outbreak occurs, so stakeholders can collaborate quickly and efficiently—be they in the U.S. or abroad; focused on humans, animals, or food; and regardless of whether they are employed by regulatory agencies, academia, or private industry.

c. Improve the coordination of emerging disease preparedness efforts among local, state, and federal officials as well as international partners. Involve key stakeholders in academia and private industry early, and often, in these planning activities.

4. Leadership and oversight

a. Determine where responsibility lies for oversight of emerging disease preparedness activities, and how such efforts will be evaluated. How will conflicting priorities be reconciled?

Implementing such an approach will face logistical and regulatory challenges. In the current regulatory system, few incentives exist to understand the complex, interconnected epidemiology of major foodborne pathogens; there is an incentive for a given agency to focus only on illnesses directly connected to a certain product falling under a certain regulatory program.

A better, more comprehensive, interdisciplinary approach is needed. Regulatory reform in particular is warranted to enable food safety agencies to monitor and address risks to the food supply beyond those occurring during slaughter or processing, in particular those originating on farms or feedlots. Here data collection and research should be focused on better understanding the complex ecology of pathogens, and on detecting emerging microbial hazards as they arise. On-farm interventions may reduce the risk of introducing new pathogens through the food system and may also have large indirect public health benefits such as lower disease transmission rates to nearby communities, lower environmental dissemination of pathogens, and lower risks of contamination to downstream produce production.

These are clearly complex goals that may not be achievable in the short or even medium term. However, concrete, incremental system-wide improvements in some of the areas mentioned above are possible within a tangible time frame and would ultimately allow the system to move toward attaining these long-term goals, even if they may not lead to immediate, radical improvements in emerging disease preparedness. To achieve these realistic but somewhat conservative changes, the U.S. should focus on the following areas:

1. Improved surveillance, specifically:

   a. Expand sampling of food animals, food production environments, and retail meats for antimicrobial-resistant bacteria, including *E. coli*, *Salmonella*, *Campylobacter*, and *Staphylococcus aureus*; these efforts, though existing, should be significantly expanded. This should include expanding NARMS sampling as well as targeted research studies. Further data are needed, particularly in the farm and feedlot environment, where it is critical to obtain a better understanding of the interplay between antimicrobial drug use, the prevalence and levels of antimicrobial-resistant pathogens in animals and the production environment, and the dissemination of these pathogens via wind, water, insects, and other pathways.
b. Continue developing improved surveillance systems (for instance, enhance whole-genome sequencing capacity), rapid detection tools, and systematic approaches for assessing vulnerabilities, both to intentional adulteration and to novel food safety hazards that so far have not been detected but that may occur in the future.

2. Improved and more timely source attribution methods, specifically:
   a. Refine methods for source attribution that will help clarify the true risks posed by pathogens such as Campylobacter, for which outbreak data seem insufficient in explaining sporadic infections. Launch new studies to inform source attribution for Campylobacter and other pathogens, possibly utilizing new subtyping methods such as MLST and whole-genome sequencing.

3. Amplified data on antimicrobial use on farms that link with surveillance data, specifically:
   a. Gather systematic on-farm data on animal drug use in cattle, poultry, and swine. This will enable better characterization of the risks of developing antimicrobial resistance among foodborne pathogens. These data should be collected and reported in a manner enabling a link to data on the prevalence and levels of antimicrobial-drug-resistant bacteria in these animals and in the foods they generate. FDA’s final rule to collect antimicrobial sales and distribution data by animal species is a promising first step in this direction.

4. Improved diagnostic methods, including rapid methods, specifically:
   a. Enhanced molecular diagnostic tools, used for future regulatory decision-making, should remain a strategic goal of the USDA. For example, the agency should develop molecular tools for identifying which STEC strains are the most hazardous and should be considered adulterants regardless of whether they belong to the “Big Six” non-O157 STECs. Focusing only on the Big Six non-O157 STECs (O26, O45, O103, O111, O121, and O145) may currently make sense, given the existing regulatory environment, current laboratory capacities, and costs, but new high-throughput molecular approaches are ultimately needed to detect other non-O157 STECs that pose important human health risks. In the meantime, important STECs that are not covered now, such as the O104 strain that caused the German outbreak, should be considered for inclusion in surveillance. New molecular tools could be based on identifying strains through genetic characteristics that have been associated with a particular public health significance—for instance, due to increases in virulence and disease severity (e.g., stx2 and eae for STEC)—or may utilize a combination of methods such as serotyping and genotyping, or may ultimately rely on whole-genome sequencing.

5. Enhanced basic research (e.g., evolution and antimicrobial resistance development), specifically:
   a. Fund further research to understand when, why, and how antimicrobial resistance emerges, and how this emergence can be predicted. Which drugs and uses pose a disproportional risk of resistance development, and which resistance mechanisms are more likely to be shared with other bacteria? In which cases will it be difficult if not impossible to reverse the emergence of resistance?

6. Strengthened epidemiological research (e.g., transmission routes), specifically:
   a. Increase research that characterizes and quantifies the foodborne risks posed by pathogens for which
foodborne transmission is unclear. Examples include *C. difficile*, multidrug-resistant *Staphylococcus aureus*, and hepatitis E virus, to name just a few. Microbiological testing and epidemiological research for these pathogens must be increased in food animals, production environments, and food products. Equally important are microbiological and epidemiological studies that better characterize and quantify human exposure risks via food. Hazard assessment models can be used to help characterize the potential foodborne risk, capture uncertainties, and target and evaluate the efficacy of intervention measures.

b. Fund additional research to investigate potential foodborne transmission to humans of important pathogens, such as drug-resistant extraintestinal pathogenic *E. coli* (ExPEC) strains, that are not typically considered foodborne.

7. Improved capabilities for rapid risk assessments that can be used for regulatory decision-making when data are scarce and conclusions are uncertain, specifically:

   a. Achieve adequate and timely assessments of potential public health risks posed by EPs and strains (for instance, new strains of *Campylobacter* spp.); this will require strong commitments to improved detection methods, surveillance (including increases in testing of animal, food, and clinical isolates), and risk assessment.

Conclusion

Just as the food system itself is not static, neither are the hazards transmitted through food. Risks may decline due to interventions or other changes in the food supply, or they may emerge or re-emerge due to factors related to the epidemiologic triangle: changes in disease agents, changes in human populations, or changes in exposure pathways. Some of these risks can be predicted better than others, but often being ready to respond quickly and effectively to an emerging microbial hazard may be more feasible than trying to predict and prevent its emergence, at least for now.

Microbial hazards in food are always changing. Humans, farm animals, wildlife, and microorganisms—both pathogenic and Nonpathogenic—live in an evolving environment and constitute a dynamic and interconnected ecosystem. Changes in food production practices or consumption patterns can result in new possibilities for the introduction, proliferation, and transmission of pathogens. Food production practices and consumption patterns change over time. New pathogens may emerge, or known pathogens may evolve to exhibit new traits. Pathogen strains may adapt to new animal reservoirs, acquire factors that increase their virulence, or acquire genes that confer resistance to antimicrobials critical to the treatment of disease in humans and animals. Demographic shifts also may determine which risks might emerge or re-emerge in the future. Aging populations are more susceptible to disease, as are young children and populations with decreased immunological status or declining overall health. In addition, our scientific understanding of the pathogens that cause foodborne infections is increasing and may cause a re-prioritization of known foodborne risks.

This report makes a number of specific recommendations. Some central themes underlie all of them: New risks will continue to emerge, and it is important to improve and integrate surveillance to be able to detect and assess
emerging hazards of both newly recognized EPs and emerging strains of known pathogens. Surveillance should span food, humans, food-producing animals, and the environments in which they live. The expeditious detection and effective control of emerging threats critically relies on the ability to recognize patterns and trends as they begin to develop; this is not possible without data that adequately capture the history and current status of the interconnected system in which we live.

**Abbreviations and acronyms**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>CDC</td>
<td>Centers for Disease Control and Prevention</td>
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<tr>
<td>EP</td>
<td>Emerging pathogen</td>
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<tr>
<td>ESBL</td>
<td>Extended-spectrum β-lactamases: an enzyme that enables microbes to resist drugs like penicillin, cephalosporins, and related beta-lactam medications.</td>
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<td>FDA</td>
<td>U.S. Food and Drug Administration</td>
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<td>FoodNet</td>
<td>Foodborne Diseases Active Surveillance Network: a CDC network that monitors foodborne disease in the United States. Established in 1995, the network includes the CDC, 10 health departments, USDA-FSIS, and FDA, with the geographic areas under surveillance covering approximately 15 percent of the U.S. population. FoodNet is an “active surveillance” network because public health officials routinely contact the more than 650 clinical laboratories within the surveillance area to make sure they appropriately report all cases to FoodNet.</td>
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<tr>
<td>USDA-FSIS</td>
<td>U.S. Department of Agriculture Food Safety and Inspection Service, an agency that regulates meat, poultry, and processed egg products.</td>
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<td>HGT</td>
<td>Horizontal gene transfer: a mechanism by which unrelated organisms share genetic information, including genetic determinants of antimicrobial resistance.</td>
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<tr>
<td>HUS</td>
<td>Hemolytic uremic syndrome: a life-threatening condition typically caused by infections with certain <em>E. coli</em> strains and characterized by the massive destruction of red blood cells, which clog the kidney’s filtration system and lead to acute kidney failure.</td>
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<tr>
<td>LEDS</td>
<td>The CDC’s Laboratory-Based Enteric Disease Surveillance, an electronic system that collects surveillance data from individual state and local public health laboratories.</td>
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<tr>
<td>MLST</td>
<td>Multi-locus sequence typing: a diagnostic method that characterizes bacteria based on their genetic makeup.</td>
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NARMS  National Antimicrobial Resistance Monitoring System: a national public health surveillance network that tracks changes in the antimicrobial susceptibility of foodborne pathogens over time by testing bacteria isolated from humans, retail meats, and food animals to determine the bacteria’s susceptibility to antimicrobial drugs.

PR/HACCP  USDA-FSIS’s Pathogen Reduction/Hazard Analysis and Critical Control Point rule, finalized in 1996, which requires meat and poultry slaughter and processing establishments to take responsibility for identifying, controlling, reducing, or eliminating foodborne hazards in their facilities.

PCR  Polynucleotide chain reaction: a method that amplifies select fragments of genetic information (e.g., one gene or part of a gene), and generates sufficient copies of the fragment to allow for sequencing (or other applications).

PPI  Proton pump inhibitors: potent antacid drugs that inhibit the secretion of gastric acid.

PulseNet  CDC’s national laboratory network that identifies foodborne outbreaks by comparing pathogens from foodborne-illness patients based on their pulsed-field gel electrophoresis (PFGE) profiles.

STECs  Shiga toxin-producing E. coli, lethal strains of E. coli that produce shiga toxins, which can severely damage organs; some of these strains can cause HUS.

Glossary of terms

Antimicrobial drug  A therapeutic drug used to treat microbial infections (e.g., antibiotics).

Antimicrobial resistance  A microbe’s ability to survive and/or grow despite the presence of a drug that would ordinarily prevent this.

Bacteriophage  A virus that infects and multiplies within a bacterium, which it can also kill.

Ceftiofur  An antimicrobial drug of the cephalosporin class (see below) used to treat bacterial infections.

Cephalosporin  A class of antimicrobial drugs that can effectively treat infections with a broad spectrum of bacteria; cephalosporins are closely related to penicillin.

Clinical isolate  The pathogen strain (or strains) isolated from clinical specimens (e.g., blood, urine, rectal swab) collected from a patient with a foodborne illness.
**Clonal strain**
A group of bacteria that share a common ancestry and are essentially identical; a clonal strain suggests the recent emergence of a new epidemic strain.

**Commensal**
A relationship between two organisms in which one partner benefits and the other is neither harmed nor benefited; commensals are normally present in the gut and on the skin.

**Culture media**
A liquid or gel that supports the growth of microorganisms or other cells in the laboratory; it is an instrumental diagnostic tool.

**Disease burden**
The impact of foodborne pathogen-related infections on public health, potentially measured as mortality rates, morbidity rates (e.g., illnesses), economic cost (e.g., treatment expenses, lost wages), or other indicators.

**Endemic**
Diseases that regularly occur in a particular population, for instance in a specific country or geographic region.

**Extra-label use**
The administration of a therapeutic drug in a way that differs from its approved use (i.e., the “label use”).

**Fluoroquinolone/Quinolone**
A class of antimicrobial drugs that can treat infections with a reasonably broad spectrum of bacteria.

**Frontline drug**
Drugs of first choice for treating human infections.

**Host range**
The range of species a pathogen can infect.

**Incidence**
A measure of disease frequency, expressed as the number of new cases within a specified time period.

**Methicillin**
An antimicrobial drug related to penicillin and used to treat penicillin-resistant infections with the bacterium Staphylococcus aureus.

**Multidrug-resistant**
A microbe’s ability to survive and grow despite the presence of multiple antimicrobial drugs, making infections from such microbes particularly challenging to treat.

**Next-generation sequencing**
A fundamentally novel method for determining genetic information, which allows the sequencing of large amounts of genetic information within a reasonable time frame.

**Nonclonal strain**
The opposite of a clonal strain (see definition above); it provides some evidence contrary to recent emergence.
**Nosocomial**
A hospital-acquired infection.

**Pathogen**
A microorganism that can cause disease (e.g., bacteria, viruses, and parasites).

**Pathogenicity**
The ability of an organism to cause disease; many microorganisms contain pathogenic and Nonpathogenic species (e.g., *E. coli*).

**Prevalence**
A measure of disease frequency expressed as the number of cases during a specified time period, including those cases with disease onset prior to the specified time interval.

**Reservoir**
The long-term “nest” of a pathogen (e.g., water, soil, wildlife) that can be the source of spillover infections into human or domestic animal populations.

**Serogroup**
Serotypes that have one or more antigens (i.e., structures to which antibodies react) in common.

**Serotype**
A strain of a microorganism as defined by serological methods; this is typically based on its surface properties, such as the presence of certain structures on a microorganism’s surface.

**Source attribution**
The process of estimating the most common food sources responsible for specific foodborne illnesses.

**Species barriers**
The mechanism that restricts a pathogen to its host range.

**Subtype**
A subset of microorganisms, usually within a serotype, that share certain properties, such as similar PFGE profiles.

**Tetracycline**
A class of antimicrobial drugs that can treat infections with a relatively broad range of bacteria.

**Trimethoprim-sulfamethoxazole**
A combination of antimicrobial drugs used to treat bacterial infections with a relatively broad range of bacterial species.

**Vector**
An organism (e.g., a tick or mosquito) that transmits pathogens from one host to another; vectors often play an important role in the epidemiology of vector-borne diseases, for example because certain stages of the pathogen can develop only inside specific vectors.
**Vehicle**

An inanimate object (also called a “fomite”) that becomes contaminated with a pathogen and transmits it to a new host (e.g., hands, clothing, trucks, and farm equipment). While vectors play an active role in disease transmission, vehicles typically play a passive one and are not required for the transmission of the pathogen.

**Zoonotic pathogen**

A pathogen that can be transmitted between animals and humans.

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### Endnotes

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22. Engering, Hogerwerf, and Slingenbergh, “Pathogen-Host-Environment Interplay and Disease Emergence.”
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31 Batz, Hoffmann, and Morris, “Ranking the Disease Burden of 14 Pathogens in Food Sources in the United States Using Attribution Data From Outbreak Investigations and Expert Elicitation.”
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Van der Waals interactions, and the like), the protein will spontaneously “fold” into some three-dimensional form that is energetically more favorable than the arrangement in a linear string. Proteins can assume more than one three-dimensional form that is somewhat favorable energetically, but proteins typically will spontaneously fold into the one specific most energetically favorable form in a given environment (e.g., a human cell). The function of proteins is closely linked to their three-dimensional form. Once folded, proteins exist in a “semi-stable” state, where proteins typically do not undergo major changes in their three-dimensional structure. Various external forces, however, such as heat, certain chemicals, or drastic changes in pH or solvent can cause proteins to temporarily or permanently change their three-dimensional form (this is, incidentally, the fundamental principle underlying “perms” and hair-straightening treatment). Such structural changes can render proteins unable to fulfill their functions.

Atypical prions are proteins that have been folded “incorrectly,” and, upon contact, can induce certain other proteins that may occur normally in the body of animals or humans to change their form and become “misfolded” themselves. The newly misfolded proteins can induce other proteins to misfold, triggering a chain reaction. These misfolded proteins can no longer fulfill their functions. In addition, the misfolded prions form aggregates (also called “fibrils”) that grow in size as more proteins are misfolded. These fibrils ultimately interfere with cell and tissue functions, causing damage to the cells and tissues in which the fibrils are formed (primarily in the central nervous system).

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