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Reassessment of the Enteropathogenicity of Mesophilic Aeromonas Species

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Cases of Aeromonas diarrhea have been described all over the world. The genus Aeromonas includes ca. 30 species, of which 10 have been isolated in association with gastroenteritis. The dominating species that account for ca. 96% of the identified strains are Aeromonas caviae, A. veronii, A. dhakensis, and A. hydrophila. However, the role of Aeromonas as a true enteropathogen has been questioned on the basis of the lack of outbreaks, the non-fulfillment of Koch’s postulates and the low numbers of acute illnesses in the only existing human challenge study. In the present study we reassess the enteropathogenicity of Aeromonas using dose response models for microbial infection and acute illness. The analysis uses the data from the human challenge study and additional data from selected outbreak investigations where the numbers exposed and the dose were reported, allowing their inclusion as “natural experiments”. In the challenge study several cases of asymptomatic shedding were found (26.3%, 15/57), however, only 3.5% (2/57) of those challenged with Aeromonas developed acute enteric symptoms (i.e., diarrhea). The “natural experiments” showed a much higher risk of illness associated with exposure to Aeromonas, even at moderate to low doses. The median dose required for 1% illness risk, was \( \sim 1.4 \times 10^4 \) times higher in the challenge study \((1.24 \times 10^4 \text{ cfu})\) compared to natural exposure events \((0.9 \text{ cfu})\). The dose response assessment presented in this study shows that the combined challenge and outbreak data are consistent with high infectivity of Aeromonas, and a wide range of susceptibility to acute enteric illness. To illustrate the outcomes, we simulate the risk associated with concentrations of Aeromonas found in different water and food matrices, indicating the disease burden potentially associated with these bacteria. In conclusion this study showed that Aeromonas is highly infectious, and that human susceptibility to illness may be high, similar to undisputed enteropathogens like Campylobacter or Salmonella.

Keywords: Aeromonas, outbreaks, water, food, challenge study, infective dose, risk of infection, risk of illness
INTRODUCTION

The genus *Aeromonas*, includes Gram-negative, non-spore-forming rods that are autochthonous of the aquatic environments worldwide (Austin et al., 1996; Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015; Graf, 2015). In humans diarrhea, and wound infections are the most common presentation followed by bacteremia (Figueras, 2005; von Graevenitz, 2007; Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). Isolation of *Aeromonas* in patients with diarrhea varies from ca 2 to 10% but a higher incidence (13%) has been found in children from Nigeria (Figueras and Beaz-Hidalgo, 2015). Also a significantly higher incidence of *Aeromonas* is found in patients with diarrhea than in those considered asymptomatic carriers (Figueras, 2005; Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). Typical susceptible patients in the different studies include babies, young children and the elderly and especially those with a pre-existing illness and/or immunocompromised (Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). According to Janda and Abbott (2010) it is important to report the presence of *Aeromonas* in stools of immunocompromised patients even if they are only asymptomatic because the risk of invasion and dissemination is considered inherently high for these patients. Nevertheless the role of *Aeromonas* in gastroenteritis has been questioned (Janda and Abbott, 1998, 2010; Chu et al., 2006; von Graevenitz, 2007), mainly because 55 out 57 challenged volunteers with a high dose of *Aeromonas* did not develop any symptoms of enteric illness, and due to the few reported outbreaks (Morgan et al., 1985; Figueras et al., 2007b; Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). However, there are also many publications providing support for a causal relation between *Aeromonas* and enteric disease (Figueras et al., 2007a,b; Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). Although 10 of the ca. 30 species that comprise the genus have been isolated in association with gastroenteritis, only 4, i.e., *Aeromonas caviae*, *A. veronii*, *A. dhakensis*, and *A. hydrophila* are the dominating species accounting for ca. 96% of the recovered isolates from this origin in different studies (Figueras and Beaz-Hidalgo, 2015).

Consumption of contaminated water and food are considered the main routes of transmission and there are few reports of well documented outbreaks that include information about the ingested doses of *Aeromonas* (Krovacek et al., 1995; Granum et al., 1998; Zhang et al., 2012). An association between diarrheal cases and consumption of untreated or contaminated drinking water has been established in several occasions (Holmberg et al., 1986; Martin Delgado et al., 2001; Khajanchi et al., 2010; Pablos et al., 2011; Ventura et al., 2015). *Aeromonas* was isolated from drinking water and stools in some patients that were diagnosed with traveler's diarrhea among the tourists that visited specific hotels in the coastal area of Tenerife and Canary Islands in Spain (Martin Delgado et al., 2001). Failures in the drinking water distribution system caused fecal contamination of the water, as revealed by the presence of bacterial indicators of fecal pollution, high concentrations of organic matter and the detection of *Aeromonas* (Martin Delgado et al., 2001). Once the water distribution system was repaired the incidence of diarrhea among the tourist population returned to normal. Even a recurrent case of *Aeromonas* bacteremia has been attributed to the consumption of contaminated well water (Katz et al., 2015). Use of contaminated water can cause secondary contamination of food products, and this can be the source of food-borne outbreaks (Altwegg et al., 1991; Krovacek et al., 1995; Granum et al., 1998; Figueras and Borrego, 2010; Wadhwa et al., 2012; Zhang et al., 2012; Figueras and Beaz-Hidalgo, 2014).

It has become generally accepted that only a subset of *Aeromonas* strains can cause gastrointestinal diseases in humans (von Graevenitz, 2007; Janda and Abbott, 2010; Grim et al., 2014). However, nowadays it is clear that infection is a complex process in which not only the virulence of the colonizing strain is important, but also its interaction with other microbes that are present in the gut, as co-infecting pathogens or in the natural microbial ecosystem, together with the specific physiological status of the host (Lund and O’Brien, 2011; Leggett et al., 2012; Beaz-Hidalgo and Figueras, 2013; Mosser et al., 2015; Ribet and Cossart, 2015; Denny et al., 2016). In fact, the overall crosstalk and interactions between commensal bacteria, enteric pathogens, and host physiology is what is considered crucial to the establishment and progression of intestinal disease (Leggett et al., 2012; Ribet and Cossart, 2015).

Challenge studies have been essential, to establish a causal relation between exposure and health effects and for quantifying the dose response relation (Teunis et al., 1996; Teunis and Havelaar, 2000). However, some outbreak reports include data on the numbers of person exposed, and even information allowing an estimate of the magnitude of the dose involved. Such outbreaks may be treated as 'natural experiments or natural exposure', comparable to a challenge experiment with a single dose group (Teunis et al., 2004, 2008, 2012b). Both data sources have been successfully used in several studies that investigated the potential of human pathogens different from *Aeromonas* for producing colonization (infectivity) and acute illness (pathogenicity) (Teunis et al., 2005, 2010).

In the present study we use dose response models for microbial infection and acute illness (Teunis et al., 1999; Teunis and Havelaar, 2000) to re-assess the Morgan et al. (1985) challenge study and combine these data with outbreak investigations to determine the dose response relation for producing infection and acute enteric disease by *Aeromonas* spp. Furthermore, to illustrate the outcomes, we estimate the risk associated with waterborne exposure to these bacteria using reference concentrations found in different water or food matrices. In addition we provide evidence in support of *Aeromonas* acting as a true enteropathogen.

MATERIALS AND METHODS

Data Used in the Dose Response Model

The volunteer protocols for the Morgan et al. (1985) human challenge study were approved by the institutional review boards of the University of Texas Health Science Center, Baylor College of Medicine, The Methodist Hospital and the General Clinical Research Center, and as common in a dose response study, did not include a non-exposed control group. The five different
Aeromonas strains that were administered orally in a bicarbonate solution to 57 adult healthy human volunteers at a range of doses and the outcomes obtained (Morgan et al., 1985) are summarized in Table 1. A prechallenge stool was cultured for enteropathogens from all volunteers before admission, and volunteers abstained from eating and drinking 90 min before and after oral challenge. Volunteers were followed up with a daily physical examination to determine symptoms of gastroenteritis (defined as 2 or more unformed stools in 24 h accompanied by any symptoms of enteric disease). Infection was defined as fecal shedding of the inoculated strain detected by analysis of the volunteer’s feces.

One of the subjects who was challenged with strain 3647 developed acute enteric symptoms, while no shedding of the inoculated strain could be detected. This outcome could indicate a false positive for symptoms, if the enteric symptoms in this subject were caused by something different, unrelated to the challenge. It could also be a false negative infection, where this subject had been colonized without detectable pathogen shedding. Such false negative shedding has been reported in other challenge studies (DuPont et al., 1969), possibly indicating intermittent shedding or low sensitivity of the detection method. Given the setting of inoculation with a high dose, we consider the second alternative (false negative infection) most plausible. In addition to these clinical experiment data, a literature review was done to collect data on acute enteric illness caused by Aeromonas due to natural exposures or outbreaks (here called “natural experiments”). Only studies where sufficient information was reported to allow a dose response assessment were selected. Four “natural experiments” with these characteristics were found: 3 foodborne outbreaks (Krovacek et al., 1995; Granum et al., 1998; Zhang et al., 2012) and an incident where a laboratory worker had accidentally ingested a pure culture of \(10^9\) Aeromonas (Carnahan et al., 1991). The data used from those studies are summarized in Table 2. To our knowledge these are the only reports that include numbers of exposed subjects and numbers of acute cases, and information on exposure, either as estimated numbers of ingested bacteria or as amount of the contaminated food product consumed (Table 2). The foodborne outbreak reported by Zhang et al. (2012) showed a dose response relation between the amounts of food consumed (cucumber salad) and the observed attack rate.

### Dose Response Assessment

Conceptually, three stages may be identified when a subject is challenged: exposure, infection, and (acute) illness. For exposure, it is important to realize that many microbial pathogens are highly infectious. Whenever a person ingests a quantity of contaminated food or drink, they only need to swallow few pathogenic particles to become infected. If the concentration of pathogens in the contaminated food or drink is low, the probability that the ingested portion did not contain any pathogens may be substantial. If (and only if) a person is exposed, there is a non-zero probability that any of the ingested pathogens survives all host barriers and succeeds in colonizing host tissues (i.e., infection is conditional on exposure). Likewise, if (and only if) a person is infected, there is a non-zero probability that the colonizing pathogen expresses pathogenicity producing damage of host tissues leading to symptoms of (acute) illness (i.e., illness is conditional on infection). Thus, there are two conditional probabilities: the probability of infection given exposure, and the probability of (acute) illness given infection. Both probabilities may be dose dependent, so that there are two dose response relations: one for infection, and the second for illness among infected subjects. Mathematical details for these two dose response relations are given in the Supplementary Material, where it is also indicated that all analyses were performed using JAGS (v4.2.0), with post-processing graphs and additional statistics in R (v3.3.1).

The parameters for these dose response relations, characterizing susceptibility to infection and to illness when infected, were estimated in a hierarchical framework. Each strain in the challenge study was treated as a separate trial, leading to strain-specific dose response relations for infection and illness. Similarly, each “natural experiment” was treated separately, as described by its own distinct dose response relations, for infection and illness. In the hierarchical framework, the variation in the parameters among these separate dose response relations is described by a (joint) distribution, representing the “group” pattern for all studies combined (Teunis et al., 2008). Hence, using all studies combined, it is possible to make predictions of the dose response relations for infection and illness, for any Aeromonas as representing a random sample from a population characterized by the combined studies shown in Tables 1 and 2.

In a subsequent analysis the challenge studies and the natural experiments were compared by assigning each of these two categories different susceptibilities to infection and illness. These separate predictions of the dose response relations for the

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**Table 1** | Strains of Aeromonas and doses challenged to different groups of volunteers (data from Morgan et al., 1985).

| Strain | Dose (cfu) | Volunteers |
|--------|------------|------------|
|        | Challenged | Infected   | Illness |
| 6Y     | 2 × 10⁴    | 4          | 1       | 0       |
|        | 1 × 10⁵    | 4          | 1       | 0       |
|        | 7 × 10⁷    | 4          | 4       | 0       |
|        | 3 × 10⁹    | 4          | 3       | 1       |
|        | 4 × 10¹⁰   | 4          | 2       | 0       |
| B158   | 6 × 10⁴    | 4          | 0       | 0       |
|        | 2 × 10⁷    | 4          | 0       | 0       |
| 3647   | 1 × 10⁷    | 4          | 1*       | 1       |
|        | 4 × 10⁷    | 4          | 0       | 0       |
|        | 2 × 10⁹    | 4          | 2       | 0       |
|        | 3 × 10¹⁰   | 4          | 1       | 0       |
| SSU    | 4 × 10⁸    | 4          | 0       | 0       |
|        | 5 × 10¹⁰   | 3          | 0       | 0       |
| 3284   | 3 × 10⁹    | 3          | 0       | 0       |
|        | 1 × 10¹⁰   | 3          | 0       | 0       |

*aAccording to Morgan et al. (1985), there was not infection but it was only acute illness, however, this could be either a false negative or a false positive (as explained in the materials and methods). Given the setting of inoculation with a high dose, we consider the first alternative (false negative infection) most plausible.*
TABLE 2 | Dose and/or concentrations of *Aeromonas* found in the ingested matrices and number of exposed and cases showing acute gastrointestinal symptoms in outbreaks or natural experiments.

| Reference            | Matrix                | Intake | Concentration cfu/g | Exposed | Ill |
|----------------------|-----------------------|--------|---------------------|---------|-----|
| Zhang et al., 2012a  | Cucumber salad        | 2 tbsp | NA                  | 178     | 73  |
|                      |                       | 1 tbsp | NA                  | 153     | 59  |
|                      |                       | 0.5 tbsp | NA              | 141     | 39  |
| Krovacek et al., 1995b | Swedish salad      | 50 g   | $10^5$–$10^7$      | 27      | 24  |
| Granum et al., 1998b | Fermented fish        | $10^7$ | 4                   | 3       |     |
| Carnahan et al., 1991b | Broth of a pure culture | $10^9$ | NA                  | NA      |     |

NA = not available; tbsp = table spoon; a Intake of food (cucumber salad) with unknown concentration, so the concentration was estimated in this study (Figure 1). b The concentration is the dose given by the authors.

RESULTS

Dose Response Assessment

The calculated estimated concentration of *Aeromonas* in the cucumber salad that was the vehicle for exposure in the outbreak reported by Zhang et al. (2012), is shown in Figure 1 and ranged between 200 and 1000 cfu/g. So the likely dose in that outbreak was lower than the doses in the challenge study (see Supplementary Material for details).

In an initial analysis the data from the challenge study and the outbreaks (natural experiments or natural exposure events) were assumed to be similar and were represented by single joint distributions (Figure 2) that showed the susceptibility to infection (i.e., the probability of infection per ingested bacteria) and to illness (i.e., the probability of illness per ingested bacteria). This unsegmented approach resulted in a single predicted dose-response relation for illness, as show in Figure 2, with a broad posterior range in illness probabilities, indicating substantial heterogeneity in illness risk. In Figure 2 it can be observed that the data from the “natural experiments” (i.e., outbreaks or natural exposure events) cluster at higher illness probabilities compared to the challenge study data. Based on these different outcomes, a refined model was set up, with separate categories for the challenge study and the “natural experiments” that represented two different distributions of susceptibility. The resulting separate dose response models for the challenge study...
and “natural exposure events” were markedly different, as shown in Figure 3. For the challenge study, ingestion of even very high doses of $10^{12}$ cfu results in low risk of acute illness (approximately 0.1, 95% range 0–0.45) as shown in Figure 3. Note, however, that even this low risk is caused by only two positive responses, with two different strains (3647 and 6Y), out of a total of 57 subjects exposed. In an alternative analysis where the symptomatic case for strain 3647 was considered a false positive, the resulting dose response models (not shown here) were virtually identical to those in Figures 2 and 3. Of those exposed, 15 were infected, indicating that infection was a lot more common than symptomatic acute illness, and also that pathogenicity may have been different among strains. For the “natural exposure events” the illness risk was much higher (approximately 0.5, 95% range 0.05–1.0), even at a comparatively low dose of $10^3$ cfu (Figure 3, lower graphic).

Figure 4 shows infection dose response relations for the challenge study and the “natural exposure” data. As infection is a condition for illness, the probability of infection must be at least as high as the illness probability (compare Figures 3 and 4). The natural exposure event data do not include any observation of infections, as this is a covert outcome (it cannot be observed directly), but because illness depends on infection the probability of infection can still be estimated. As the illness risk was high in these “natural experiments”, the infection probabilities must be high as well, as shown in Figure 4.

The estimated susceptibilities for infection and illness were illustrated also by the doses required to cause a 1% probability of infection for each of the challenge strains, and separately for each of the “natural exposure events” (Table 3; Figure 5). Estimates for the two prediction categories: low susceptibility for the challenge studies and high susceptibility for the “natural experiments” are shown in Figure 5 for infection and for illness risks. An outcome of 1% risk was chosen here because in the challenge studies the risk was so low that calculation of 50% infectious (and illness) doses results in unrealistically high numbers. Mean and median, as well as a 95% range are also given in Table 3.

**Estimation of Risk**
To illustrate the application of the dose response models derived here, a small risk study was set up, using a simple scenario...
of three different, fixed doses, representing low, medium, and high exposure. **Figure 6** shows the resulting risks of *Aeromonas* infection and illness. It is clear that using the challenge study dose response relation, the illness risk remains small, even for exposure to considerable doses of *Aeromonas*. On the other hand, the natural exposure events dose response relation produces high illness risks, even when the dose is moderate or low.

**DISCUSSION**

The major clinical manifestation produced by *Aeromonas* is diarrhea affecting principally young children and immunocompromised patients (Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). The single available human challenge study for *Aeromonas* shows low susceptibility, in particular to developing acute enteric symptoms with only 2/57 (3.5%) challenged being positive (Table 1; Morgan et al., 1985). In contrast, the “natural exposure events” indicate that a high risk of illness may be associated with exposure to *Aeromonas* even at moderate to low doses. Thus, host susceptibility may be a strong determinant for the illness risk of *Aeromonas* sp. The dose response assessment presented in this study corroborates this conclusion (median 1% illness dose was $\sim 1.4 \times 10^4$ times higher in “the natural exposure events” compared to the clinical challenge), and shows how these two subsets of illness data lead to different estimates of susceptibility.

Although invaluable for understanding infection and pathogenesis, clinical challenge may not quantitatively represent natural infection that occur during an outbreak. Clinical challenge may underestimate the risk because pathogen inocula may decrease in virulence because of safety testing, often requiring repeated culture in the laboratory. Likewise, volunteers are selected for being healthy and immunocompetent to reduce the risk of serious complications. Conversely, outbreaks select for virulent pathogen strains and susceptible hosts, as this increases the probability that a cluster of cases is detected (Teunis et al., 2004). Consequently, challenge studies and outbreaks may both be biased, in opposite directions: each may represent an extreme in the continuum of dose response relations as discussed earlier (Teunis et al., 2005, 2010; Thebault et al., 2013). Outbreaks or natural exposure events may select for highest susceptibility in the affected host population producing a high illness risks, even when the dose is moderate or low. For instance in the food outbreak that produced acute diarrhea in over 200 college students’ described by Zhang et al. (2012) the estimated concentration of *Aeromonas* in the cucumber salad determined in the present study ranged from 200 to 1000 cfu/g (Figure 1). This estimated concentration assumed a homogeneous distribution of bacteria (Teunis et al., 2005) in the cucumber salad. As the origin of the contamination in this outbreak was attributed to rinse water of the vegetables, contaminated with sewage, occurrence of bacteria may not have been severely clustered. In case there is severe clustering, some subjects would have been exposed to high doses, while others would not have been exposed at all. Only when there is extreme clustering, the slope of the dose response relation is affected, not its location and the mean dose remains proportional to the amount of contaminated food that was ingested (Teunis et al., 2008, 2012a). As no information regarding clustering in the contaminated cucumber salad was provided by Zhang et al. (2012), any assumptions regarding clustering would be speculation.

Little is known about who is susceptible to any specific strain or pathogen in the population at large, or how many of the already considered susceptible like infants or immunocompromised would develop acute illness. It has been estimated that in developed countries the part of the population that could be susceptible to acquiring a foodborne disease would be near 20% (Lund and O’Brien, 2011; Lund, 2015). Challenge
Giardia lamblia (26%) were infected. The occurrence of few illness outcomes is not a true gastrointestinal pathogen (von Graevenitz, 2007; Janda et al. 2010). Studies directed by Dr. Chopra, i.e., Grim et al., 2014 and Beaz-Hidalgo, 2015, found in several studies (Svenungsson et al., 2000; Vila et al., 2003; Figueras, 2005; Senderovich et al., 2012; Figueras and Beaz-Hidalgo, 2015). In fact even 30% of the genomes deposited at the GenBank database with the name of *A. hydrophila* do not belong to this species (Figueras et al., 2014; Beaz-Hidalgo et al., 2015). Old literature on clinical aeromonads limited the identification only to three species, i.e., *A. hydrophila*, *A. sobria* (the correct terminology for the clinical strains is *A. veronii* biovar sobria) and *A. caviae* (Janda and Abbott, 1998; Figueras, 2005; von Graevenitz, 2007). However, strains under these names may belong to other species (Janda and Abbott, 1998; Soler et al., 2003; Figueras, 2005). For instance in the Morgan et al. (1985) study among the five strains named *A. hydrophila* three (6Y, B158, and 3284) were originally thought to belong to *A. sobria* (the correct terminology *A. veronii* biovar sobria), and strain SSU, as commented above, corresponds to *A. dhakensis*, a highly virulent species (Beaz-Hidalgo et al., 2013; Morinaga et al., 2013; Figueras and Beaz-Hidalgo, 2015), instead of *A. hydrophila* as originally thought (Morgan et al., 1985). It is now known that the importance attributed to *A. hydrophila* is due to misidentification of the majority of strains as belonging to this species, using phenotypic identification systems (Soler et al., 2003; Figueras, 2005; Beaz-Hidalgo et al., 2010; Morinaga et al., 2013). In fact even 30% of the genomes deposited at the GenBank database with the name of *A. hydrophila* do not belong to this species (Figueras et al., 2014; Beaz-Hidalgo et al., 2015). The above analysis suggests that the natural exposures involve much lower doses than the challenge study. Notwithstanding the low risk predicted from the challenge study data, the same study suggests that the risk of infection is not extremely low, in particular for two of the strains 6Y and 3647 (Morgan et al., 1985). This means that the risk cannot be ignored, even though few cases of acute diarrhea were observed. Janda and Abbott (2010) underlined that the strain SSU (CDC diarrheal isolate), which is probably the most well-characterized *Aeromonas* strain with respect to virulence factors (i.e., carriage and expression of enterotoxin genes, and potential colonization factors see studies directed by Dr. Chopra, i.e., Grim et al., 2014 and Ponnusamy et al., 2016, and references therein) did not produce infection (colonization) nor diarrhea in the challenge study, so this cannot support the idea that the strains were wrongly selected. However, it is still possible that a critical virulence or colonization factor could have been lost when subculturing the original strains (Morgan et al., 1985; Janda and Abbott, 2010). Experimental studies in a murine animal model showed that the injection of strain SSU generated the death of all mice in 48 h (Grim et al., 2014; Ponnusamy et al., 2016). Strain SSU has been recently identified as *Aeromonas dhakensis*, a highly virulent species (Beaz-Hidalgo et al., 2013; Morinaga et al., 2013; Figueras and Beaz-Hidalgo, 2015), instead of *A. hydrophila* as originally thought (Morgan et al., 1985). It is now known that the importance attributed to *A. hydrophila* is due to misidentification of the majority of strains as belonging to this species, using phenotypic identification systems (Soler et al., 2003; Figueras, 2005; Beaz-Hidalgo et al., 2010; Morinaga et al., 2013). In fact even 30% of the genomes deposited at the GenBank database with the name of *A. hydrophila* do not belong to this species (Figueras et al., 2014; Beaz-Hidalgo et al., 2015). Strain SSU is due to misidentification of the majority of strains as belonging to this species, using phenotypic identification systems (Soler et al., 2003; Figueras, 2005; Beaz-Hidalgo et al., 2010; Morinaga et al., 2013). In fact even 30% of the genomes deposited at the GenBank database with the name of *A. hydrophila* do not belong to this species (Figueras et al., 2014; Beaz-Hidalgo et al., 2015). Strain SSU is due to misidentification of the majority of strains as belonging to this species, using phenotypic identification systems (Soler et al., 2003; Figueras, 2005; Beaz-Hidalgo et al., 2010; Morinaga et al., 2013). In fact even 30% of the genomes deposited at the GenBank database with the name of *A. hydrophila* do not belong to this species (Figueras et al., 2014; Beaz-Hidalgo et al., 2015).
FIGURE 5 | Box plots showing the estimated doses required for 1% probability of infection (InfD01, upper graphic) and illness (IllD01, lower graphic). Results in both graphics are provided for each individual strain (6Y, B158, 3647, SSU, and 3284) and experiment (Zh = Zhang et al., Kr = Krovaček et al., Gr = Granum et al., Ca = Carnahan et al.) and globally for all strains (i.e., low susceptibility) and natural experiments (i.e., high susceptibility).

also in previous reviews (von Graevenitz, 2007). However, studies on traveler's diarrhea where an adult population is involved are dominated by *A. veronii* (Vila et al., 2003; von Graevenitz, 2007 and references therein). Also in a recent study performed by Chen et al. (2015) in and adult population from Taiwan *A. veronii* was the prevailing species (54.6%) followed by *A. caviae* (27.6%), and with one strain of each of the species *A. dhakensis* (9.1%) and *A. sanarellii* (9.1%). Interestingly one patient in the latter study had a history of eating lettuce with salad prior to the illness. In a recent study tomatoes and parsley irrigated with the same water showed the same genotype of a strain of *A. sanarellii* and the same occurred for an *A. caviae* strain recovered from the water and the irrigated lettuces (Latif-Eugenín, 2015).

As we commented in the introduction it is generally assumed that the virulence of *Aeromonas* is multifactorial and that only a subset of *Aeromonas* strains is capable of causing gastroenteritis in humans (von Graevenitz, 2007; Janda and Abbott, 2010; Beaz-Hidalgo and Figueras, 2013; Figueras and Beaz-Hidalgo, 2014; Grim et al., 2014; Ponnusamy et al., 2016). In fact some *Aeromonas* strains possess several virulence toxins and secretion systems, among which the Type III secretion system and the Shiga toxin genes are similar to those present in other important pathogenic bacteria (Figueras et al., 2007a; Alperi and Figueras, 2010; Figueras and Beaz-Hidalgo, 2014). The large variation observed among *Aeromonas* strains maybe linked to the site where the virulence is expressed (Leggett et al., 2012). For instance carriage of virulence genes that have a local action within the host cells like, i.e., the injection of toxins through the Type III or Type IV secretion systems may generate a more virulent response than other strains that exert virulence at more distant sites (Leggett et al., 2012). The latter occurs when pathogens cause secretion of proteins binding to host cells (i.e., immune modulators delivered by the general secretary pathway, or by Type I, II, and V secretory systems). This different pathway or the synergetic effect of such pathways may lead to differences in infective doses (Leggett et al., 2012). In addition the host susceptibility is very relevant, now we know that the genetic polymorphism in the host population determines the variability observed in the type or intensity of responses against the encountered specific pathogens ranging from asymptomatic infections to fatal disease (Ribet and Cossart, 2015).

The concentrations of *Aeromonas* that were selected in the exposure scenarios to assess risk of infection, were those that can normally be found in oysters, i.e, MPN 9.3/100 g or 10 cfu/100 ml have been reported in several drinking water studies (Abeyta et al., 1986; Austin et al., 1996; Figueras et al., 2005; Egorov et al., 2011; Robertson et al., 2014) or for instance in Milk ($1 - 2 \times 10^3$) or meat products ($10^2 - 10^3$) as found by Borrell et al. (1998).
In fact a MPN 9.3/100g was the concentration of *Aeromonas* found in oysters considered to be the source of an outbreak that affected 472 persons suffering from gastroenteritis in Louisiana (Abeyta et al., 1986). Concentrations as high as $10^5$ or $10^6$ cfu/100 ml can be found in treated wastewater and in reclaimed waters used for irrigation of vegetables (Austin et al., 1996; McMahon and Wilson, 2001; Janda and Abbott, 2010; Fernandez-Cassi et al., 2016). Such sources could contaminate ready to eat vegetables and pose a risk for human health as was recently demonstrated finding the same genotype of *Aeromonas* in the irrigated water and in the irrigated vegetables as commented above (Latif-Eugenín, 2015).

The dose response assessment reported here shows that when *Aeromonas* is present, the probability of infection may not be negligible, and in a susceptible host, there may be a high risk of acute enteric illness. It is therefore important to determine how rare it is for a host to be highly susceptible. Outbreaks or other natural experiments are hard to find. This may be because they are so rare, but there could be underreporting because *Aeromonas* is often not considered among the microbes to be analyzed during investigation of outbreaks of infectious gastroenteritis (Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). In fact the recognition of *Aeromonas* in the clinical setting occurs accidentally on routine enteric isolation media designed for other enteropathogens like XLD (MacConkey, Xylose Lysine Dextrose Agar), SS (Salmonella–Shigella Agar), or CIN (Cefsulodin–Irgas–Novobiocin Agar) as described earlier (Figueras, 2005; Janda and Abbott, 1998, 2010; Figueras and Beaz-Hidalgo, 2015).

The carriage of an infectious microorganism in the general population cannot be easily extrapolated from (observable) symptomatic cases, because of underreporting, but also because in the population at large, a great majority of the infections may remain asymptomatic. Biomarker studies using serum antibodies for estimating infection (seroconversion) rates in the general population have shown that for *Campylobacter* there may be more than 1,000 asymptomatic infections for every notified case of campylobacteriosis (Teunis et al., 2012b, 2013). Similar ratios have been found for *Salmonella* (Simonsen et al., 2011). Serology could be helpful in determining whether there could be substantial carriage of *Aeromonas* in the general population. Serological evidence of infection has been provided in some cases, supporting the true enteropathogenicity of *Aeromonas* (see review by Janda and Abbott, 1998; Crivelli et al., 2001). In fact a specific secretory immunoglobulin A (sIgA) response at the intestinal mucosa against the extracellular products that appeared in the feces of patients with *Aeromonas* diarrhea was demonstrated by Crivelli et al. (2001).

Prevalence or incidence of HIV, hepatitis C have been estimate using the “evidence synthesis”, that is a well-accepted methodology for integrating various sources of data to estimate a quantity of interest for which there are no or limited direct data (McDonald et al., 2015 and references therein). This approach was applied recently to determine the incidence of symptomatic pertussis infection in the Netherlands (McDonald et al., 2015) and may help in determining whether the low numbers of symptomatic cases are consistent with frequent isolation of *Aeromonas*.

The predicted risks of infection and illness show considerable uncertainty. This is caused by the variation observed between outcomes produced by different strains, but it also results from the small sizes of the exposed groups, in particular in the challenge study (Table 1). Unfortunately it is not likely that there will be more or better data available, to improve the precision of the estimates. Despite the small sample sizes and uncertain doses, what is relevant is that the low and high susceptibility estimates are clearly different.

**Aeromonas is a True Enteropathogen**

Arguments used against considering *Aeromonas* an enteropathogen, i.e., the lack of outbreaks, the non-fulfillment of the Koch’s postulates, the low numbers of acute illness in the human challenge study and the lack of animal model have all been addressed with evidence to the contrary in other studies (Chu et al., 2006; Figueras et al., 2007b; von Graevenitz, 2007; Janda and Abbott, 2010; Figueras and Beaz-Hidalgo, 2015). A murine model of *Aeromonas* diarrhea has been developed (Abuelsaad et al., 2013). The Koch’s postulates have been fulfilled by considering the incidental ingestion of *A. trota* by a...
laboratory worker (Carnahan et al., 1991), the challenge study of Morgan et al. (1985) and the outbreak reports included in this study (Krovaček et al., 1995; Granum et al., 1998; Zhang et al., 2012). In addition, even though few outbreaks exist, an epidemiological link has been found between the source of infection and the clinical isolates. The same Aeromonas strains (verified by genotyping) that caused diarrhea were isolated from drinking water (Khajanchi et al., 2010; Pablos et al., 2011), from the consumed shrimp cocktail (Altweg et al., 1991) and from the household environment (Demarta et al., 2000). Also the same genotype of Aeromonas has been isolated from an HIV/AIDS patient suffering from gastroenteritis and from their household drinking-water (Ramalivhana et al., 2010).

CONCLUDING REMARKS AND PERSPECTIVES

Combined evidence collated from clinical studies in humans and outbreaks shows that Aeromonas should be treated as a human enteropathogen. Exposure to low doses of Aeromonas sp. may lead to infection, but most infections may remain asymptomatic. Given the omnipresence of Aeromonas in the environment, seroprevalence studies in the general population are needed to reveal frequent carriage. Further investigations are needed to determine the specific combination of host, environment and pathogen factors that lead to the occurrence of acute enteric symptoms (illness) associated with Aeromonas infections. The risk of illness may be considerable, even when exposed to moderate doses as shown for the studied outbreaks. Therefore, as suggested earlier, patients with underlying malignancies or immunosuppressing conditions should be closely supervised, considering the inherently high risk of invasion and dissemination associated with this population group.

AUTHOR CONTRIBUTIONS

PT and MF contributed equally to the conception and design of the study, PT performed the mathematical modeling and PT and MF jointly interpreted the data and wrote the manuscript. Both authors approved the final version of the paper.

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SUPPLEMENTARY MATERIAL

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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