**Toxoplasma gondii** in beef consumed in France: regional variation in seroprevalence and parasite isolation

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**Abstract** – In France, the consumption of cattle and sheep meat appears to be a risk factor for infection of pregnant women with *Toxoplasma gondii*. Several nation-wide surveys in France have investigated the prevalence of *T. gondii* in sheep and pig meat, but little is known at present about the prevalence of the parasite in beef. The main objective of the present cross-sectional survey was to estimate the seroprevalence of *T. gondii* infection in beef consumed in France. A secondary objective was to attempt to isolate *T. gondii* from cattle tissues and to study the geographical and age variations of this seroprevalence. The overall estimate of seroprevalence of *T. gondii* in bovine carcasses (*n* = 2912), for a threshold of 1:6 was 17.38%. A strong age effect was observed (*p* < 0.0001) with a seroprevalence of 5.34% for calves (<8 months) and 23.12% for adults (>8 months). Seroprevalence estimates given by area of birth and area of slaughtering for adults showed that the areas with the highest seroprevalence were not the same between these two variables. Only two strains, corresponding to genotype II, were isolated from heart samples, indicating that there is a limited risk of human infection with *T. gondii*, which needs to be correlated with the food habit of consuming raw or undercooked (*bleu* or *saignant*) beef. However, new questions have emerged, especially concerning the isolation of parasites from beef and the precise role of bovines, generally described as poor hosts for *T. gondii*, in human infection.

**Key words:** *Toxoplasma gondii*, Bovines, Meat, Strain isolation, Prevalences.

**Résumé** – *Toxoplasma gondii* dans la viande bovine consommée en France : variation régionale de la séroprévalence et isolalement de parasites. En France, la consommation de viande bovine et ovine apparaît comme un facteur de risque pour la contamination des femmes enceintes par *Toxoplasma gondii*. Plusieurs enquêtes nationales ont été réalisées afin de déterminer le niveau de contamination par *T. gondii* de la viande ovine et porcine, en France, mais très peu est encore connu quant à la prévalence du parasite dans la viande bovine. La présente enquête transversale avait pour objectif principal d’estimer la séroprévalence de l’infection à *T. gondii* dans la viande bovine consommée en France, ainsi que d’isoler *T. gondii* à partir de tissus de bovins et d’étudier, à titre d’objectif secondaire, les variations géographiques et d’âge de cette prévalence. L’estimation globale de la séroprévalence de *T. gondii* dans les carcasses de bovins (*n* = 2912) était de 17.38 % (pour un seuil de dilution à 1:6). Un effet significatif de l’âge a été observé (*p* < 0.0001) avec une séroprévalence de 5.34 % pour les veaux (<8 mois) et de 23.12 % pour les adultes (>8 mois). Les estimations de séroprévalence données par zone de naissance et par zone d’abattage pour les parasites de *T. gondii* indiquent un faible risque pour la contamination des femmes enceintes.

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Introduction

Toxoplasma gondii is an obligate intracellular parasite belonging to the Apicomplexa phylum, with world-wide distribution. It has three infectious stages: the tachyzoites, the bradyzoites, and sporulated oocysts (with eight sporozoites). These stages are linked in a complex life cycle, with the cat belonging to the Apicomplexa phylum, with world-wide distribution. All warm-blooded species, including humans, can act as intermediate hosts [53]. The main route of transmission is oral: humans and animals can become infected by ingesting either tissue cysts containing bradyzoites, from raw or undercooked meat, or sporulated oocysts, from various environmental sources (soil, water, vegetables). Consumption of undercooked meat may be considered a major risk factor for humans, especially in Europe, where it has been associated with various sources of infection, four to sixty-three percent of infections [14, 56], while Hill [35] has demonstrated the predominance of oocyst-driven infections in North America. In humans, infection is usually either asymptomatic or the cause of mild flu-like symptoms. However, toxoplasmosis can be life-threatening especially in immunocompromised individuals. Moreover, if acquired during pregnancy, toxoplasmosis can cause miscarriage or congenital malformations affecting the brain, eyes or other organs of the foetus [34]. In animals, infection with T. gondii may cause abortion in sheep, goats and pigs [28], while in cattle, natural T. gondii infection does not appear to cause clinical disease or abortion [23]. Therefore, interest in T. gondii in cattle comes mainly from a public health perspective: if cattle carry viable tissue cysts, they may be a source of human infections. This might be particularly true in western countries, where eating raw or undercooked beef, is to some extent, related to high standards of living [28]. Moreover, several human toxoplasmosis outbreaks were recorded world-wide, with various sources of infection, four of them identifying the consumption of raw or undercooked beef as the most probable source [2, 54]. However, the quantified role of these animals in the transmission of T. gondii to humans remains unclear. On the one hand, serological evidence of T. gondii exposure in cattle has been published world-wide [2, 10, 28, 31, 36–41, 45, 47, 48, 50, 51, 55, 57], and parasite DNA has been detected by means of polymerase chain reaction (PCR) [5, 32, 49, 50]. On the other hand, a few attempts have succeeded in isolating the parasite from various types of bovine tissues, by means of bioassays [7, 23, 25, 50, 51].

In spite of a constant decrease in the last 40 years, France is still the leading consumer of beef within EU with 24 kg/inhabitant in 2013 [30]. When it comes to cooking habits, a large majority (>50%) of French adults consume undercooked (bleu, saignant, à point) lamb, beef or horse meat, while 20% of them eat, at least occasionally, raw beef [2]. Not surprisingly, the consumption of cattle and sheep meat has appeared as a risk factor for infection of pregnant women in France [6]. Similarly in Italy, in a recently published quantitative risk assessment model to estimate the yearly possibility of acquiring human toxoplasmosis infection due to consumption of beef or pork, beef scored higher [8]. Few previous studies [2, 31] have investigated T. gondii infection in cattle in France, while a nation-wide survey was conducted to evaluate the prevalence of T. gondii in sheep and pig meat consumed in France [19, 33]. Since several studies have pointed out the need for large-scale screening to clarify the role of cattle as carriers of viable tissue cysts [49, 50], in the present study, we used a cross-sectional survey to estimate, as a primary objective, the seroprevalence of T. gondii infection in beef consumed in France. As a secondary objective, we attempted the isolation of T. gondii from cattle tissues and studied the geographical and age variations of this seroprevalence.

Materials and methods

Sampling strategy

Sampling size

For an expected seroprevalence ranging from 5% to 43%, as previously observed in different studies in cattle [28, 31, 50, 57], a sample size of 3000 animals enabled us to guarantee a relative precision of minimum 0.2% [20]. Two different types of meat were targeted in terms of origin: national production (slaughtered in France) and imports (slaughtered within the EU or outside the EU). The relative portion of imports of fresh beef was estimated in 2007 to be 15% and 20% (Data of the French Ministry of Agriculture). The number of samples from national production was set originally at 2350 and from imports at 650.

Samples of French origin

The sampling strategy was established in order to be representative of the spatial variability of slaughterhouses and the age of animals. The first unit of sampling was the slaughterhouse. From French Ministry of Agriculture databases for 2007, 245 cattle slaughterhouses were identified. The second unit of sampling was 50 animals in a single week from the same slaughterhouse for logistical reasons. In order to obtain enough samples from the 245 slaughterhouses, we first excluded 111 with less than 5200 animals/year (less than 100 per week), which represented 4.66% of production. We also excluded two overseas slaughterhouses (to limit costs). The 25 largest slaughterhouses were included a priori in the sampling, representing 50.94% of meat production. We selected a minimum of one slaughterhouse per administrative region.
(n = 21), with probability of sampling proportional to meat production (unequal probability of sampling) using the SURVEYSELECT procedure in SAS. One slaughterhouse was added to the sampling protocol and selected among three other slaughterhouses in the Pyrénées-Atlantiques department because of the high seroprevalence rate for ovine *T. gondii* infection, as shown in our previous results [33]. Therefore 47 different slaughterhouses were selected at the beginning of the study, but due to practical reasons, 48 were included (one region performed sampling in two different slaughterhouses), covering all regions of France (Fig. 1). It was also checked that the age distribution of the sampled animals was similar to that of cattle slaughtered at the national level. The percentage of calves (<8 months) was fixed, taking into account the yearly proportion of veal production in each slaughterhouse. The percentage of calves in the sample was set at the national level at 25%, bearing in mind that it was estimated at around 30% of French production in heads at slaughterhouses in 2007 (around 15% in tons).

**Imported samples**

Sampling was performed at the International Food Market (IFM) of Rungis (Ile-de-France [IDF] region), which is the largest food market for trading (national/international) of agro-food products (fruits, vegetables, dairy and other animal products, seafood, etc.) in France. The sampling plan enabled a representative allocation between the sources of imported carcasses, while covering 99% of the origins of the bovines, with a minimum of 30 analyses per country of origin.

**Sampling protocol**

All samples were collected between January and July 2009. For the samples of French origin, only the heart was collected, while for the imported carcasses only the diaphragm was collected. An animal less than 8 months old was considered a calf, while an animal older than 8 months was considered an adult.

**Samples of French origin**

Fifty fresh hearts were collected per slaughterhouse and two slaughterhouses were sampled weekly. In the selected slaughterhouses, systematic random sampling was applied. The pace of sampling between two carcasses was defined by the ratio of the number of animals to be slaughtered that week/50 (animals to be sampled in 1 week). This method was designed to reduce the over-representation of animals raised together [13]. Hearts were collected in individual plastic boxes and kept refrigerated (+4 °C) for maximum 3 days before being sent to the National Reference Centre for toxoplasmosis in Reims (NRC, Reims, France) or to the National Reference Laboratory for food-borne parasites in Maisons-Alfort (NRL, Maisons-Alfort, France). Each laboratory received 50 fresh heart samples weekly, originating from only one slaughterhouse, according to the schedule plan.

Data about age and birth origin were collected for each sampled animal.

**Imported samples**

No hearts were available for the imported bovines, since they arrive in France as eviscerated carcasses. Therefore, the diaphragm was the only available sample, collected monthly and stored at −20 °C before being sent to the NRL (Maisons-Alfort, France).

**Serological tests**

The modified agglutination test (MAT) for the detection of *T. gondii*-specific immunoglobulin (immunoglobulin G) antibodies was performed, as previously described, on all cardiac and diaphragm fluids using an antigen prepared from formalin-fixed whole RH tachyzoites produced and quality-controlled
by the NRC (Reims, France) [33]. Cardiac fluids were collected by pipetting the exudate on the bottom of each plastic box. Meat juice was obtained from 25 g of diaphragms cut into small pieces and frozen overnight at −20 °C in a plastic bag. After thawing at room temperature, the meat juice was collected with a pipette into a microtube, as previously described [46]. The fluids were serially two-fold diluted. The threshold dilution was 1:6.

**Bioassay for *T. gondii* in mice**

The *in vivo* experiments were approved by the local Animal Research Ethics Committee (ComEth Ansès, EnvA, UPE) of Maisons-Alfort. In order to limit suffering and distress, mice were acclimated for 7 days after their arrival. Cages were filled with paper strips. Animal health and behaviour were monitored daily. Mice were observed based on the following criteria:

- external physical appearance (disheveled or spiked hairs, watering eyes, bent back, tremors),
- behaviour (exploratory behaviour decrease, unusual posture, prostration),
- behavioural response to external stimuli (no response).

If any of these criteria were critically altered, mice were subsequently euthanised and examined post-mortem to investigate the parasitic load. Euthanasia consisted in CO₂ asphyxiation, followed by cervical dislocation.

After approval from the local Animal Research Ethics Committee, between 5 and 11 heart samples, randomly chosen from among those with the highest titers of agglutinating antibodies, were bioassayed weekly in three outbred female Swiss Webster mice (Charles River Laboratory, France). Additionally, nine seronegative hearts, randomly selected from the total number of samples, were bioassayed in mice. Briefly, each whole heart was mixed and incubated at 37 °C for 1.5 h with trypsin (final concentration 0.25%). The suspension was then filtered, pelleted by centrifugation, washed in saline, and resuspended in a saline solution containing penicillin G, streptomycin, and amoxicillin to limit bacterial proliferation. This homogenate was inoculated intraperitoneally into three mice [1, 59]. Mice were monitored twice daily with food and water supplied *ad libitum*. In case of acute toxoplasmosis, the mice were culled by CO₂ asphyxiation, followed by cervical dislocation, and samples of brain were taken for analysis. Mice were bled 4 weeks post-inoculation and their serum was tested at 1:25 dilution for *T. gondii* antibodies with the MAT. Later on, mice were culled 60 days post-inoculation by cervical dislocation, and their brains were examined for tissue cysts.

**Genotyping of *T. gondii* isolates**

Brain cysts from seropositive mice were isolated by percoll gradient centrifugation [15]. DNA was extracted using a QIAGen DNA MiniKit (Qiagen, Courtaboeuf, France), and genotyping analysis of *T. gondii* DNA was performed with 15 microsatellite markers in a single multiplex PCR assay, as described elsewhere [3]. All strains isolated were cell cultivated and banked in the *Toxoplasma* Biological Resource Center (BRC, Reims).

**Statistical analysis**

All analyses were performed in R (version 3.5.1), using the survey package (version 3.34) to fit the complex survey sample [42, 43]. Individual data were weighted for unequal sampling probability. Post-stratification was made in order to have two slaughterhouses and a minimum of 100 animals by geographical area. The complex design of the survey, considering the slaughterhouse as a cluster, and the geographical area as strata was taken into account in the statistical analysis and seroprevalence estimates. Imports were assumed to represent 15% of production. Imports were considered in French national seroprevalence estimates of beef, even though the tissue used for analysis was different (diaphragm), in order to obtain an overall estimate of exposure in France. Estimates of the differences in sensitivity or specificity between the two tissues (heart/diaphragm) used for autochthonous samples or imports were not available at the time of analysis, and as a consequence, these differences were not taken into account. Descriptive analyses were done for different dilutions in MAT at 1:6 to 1:200.

For French production, we analysed the probability of being seropositive with different explanatory variables such as the influence of age, sex, type of production (milk or meat), area of birth, life and slaughtering. Univariate analyses for the variable “age” (in the category “adult” or calf) and the variable “origin” (in the category “import” or “France”) were done using the Wald test.

For multivariate analysis, a sample showing a positive result in a 1:6 dilution in MAT was considered seropositive. The serological status was coded as 1 (positive) or 0 (negative). The response variable was analysed using logistic regression (quasi-binomial distribution and logit link function). The effect of age in years was also tested as a quantitative variable, with different polynomial degrees from linear regression to sixth degree polynomial regression. A first order interaction term was also tested. All possible models were compared using the lowest Akaike Information Criteria (AIC) [11]; AIC is defined using Rao-Scott approximation [44]. When several models show a difference in AIC less than 2, the model with the fewest number of parameters was retained, the most parsimonious one. The significance of each effect was determined using the Wald test.

**Results**

**Sample collection**

**Samples of French origin**

All 48 selected abattoirs were part of the sampling plan, except one (Vitré, Bretagne region). Therefore, two-fold increased sampling was carried out in one abattoir (Liffére, Bretagne region). A total of 2349 samples were collected; 574 of them were from calves and 1774 were from adults. For one sample, a qualitative indication of age (adult/calf) was not available. The area of birth was known for 1990
samples (359 unknown), of which 1473 were adults and 517 were calves. The minimum age at slaughtering in the dataset is 0.2 years, and the oldest animal was 20.4 years old. The mean age at slaughtering (taking into account the design of the survey), is 3.2 years, for a calf 5 and a half months (0.47 years), and for an adult 4.35 years old. Around 3% of observations (79/2349) were above or equal to 12 years old.

Imported samples

A total of 650 diaphragms were collected at the International Food Market (IFM) of Rungis (Ile-de-France (IDF) region). For 79 of them, no muscle juice was obtained due to an advanced condition of dehydration of the meat. Another seven were missing the origin country data. One was collected as an imported sample, but was in fact of French origin. A total of 563 animals were thus included in the statistical analysis (imports included) was found to be 17.38% [12.73–23.12] (Table 2), with a specificity of 90.7% of the relative frequency in French meat imports (Table 7). Moreover, 225 samples were from calves and 338 from adults, without a precise indication of the age (in years or months).

Serological tests

The results of the MAT according to the origin of samples (French/imported), age of animals (calves/adults), and terminal titer are presented in Table 1. In total, there were 318, 124, 117, 16, 12, 10 and 1 sample positive at a terminal titer of 6, 10, 25, 50, 100, 200 and 400, respectively.

Seroprevalence

Overall seroprevalence

The overall seroprevalence in beef consumed in France (imports included) was found to be 17.38% [12.73–23.26] at a cut-off titer of 6. A decrease was observed when a higher cut-off was defined: at a titer of 10, the overall seroprevalence was 8.93 [5.97–13.17] and at a titer of 25, the overall seroprevalence was 5.01 [2.73–9.02] (Table 2).

The seroprevalence for calves was estimated to be 5.34% [2.49–11.09] at a cut-off titer of 6 (Table 3), and for adults 23.12% [17.83–29.40] at a cut-off titer of 6 (Table 4). As found previously, the seroprevalence decreased for both categories when a higher cut-off was chosen. Age as a qualitative variable (calves/adults) was found to be highly significant in univariate analysis ($p$ significance or $p$-value = $2 \times 10^{-5}$).

Samples of French origin

The overall seroprevalence of *Toxoplasma gondii* in cattle slaughtered in France was estimated at a cut-off titer of 6 to be 16.96% [11.91–23.56] (Table 2), with a specific seroprevalence of 5.06% [1.79–13.50%] for calves (Table 3) and 22.00% [16.16–29.22%] for adults (Table 4). As found previously, the seroprevalence decreased for both categories, when a higher cut-off was chosen.

### Table 1. Results of the modified agglutination test (MAT) accordingly to the origin of samples (French/imported), age of animals (calves/adults) and terminal titer.

| Number of examined samples | Number of samples with terminal titer of: |
|----------------------------|------------------------------------------|
|                            | 0  | 6  | 10 | 25 | 50  | 100 | 200 | 400 |
| Samples of French origin   |    |    |    |    |     |    |     |     |
| Calves                     | 574| 535| 25 | 6  | 4   | 3   | 1   | 0   |
| Adults                     | 1774| 1334| 266| 90 | 52  | 12  | 11  | 8   |
| Imported samples           |    |    |    |    |     |    |     |     |
| Calves                     | 225| 211| 3  | 3  | 8   | 0   | 0   | 0   |
| Adults                     | 338| 233| 24 | 25 | 53  | 1   | 0   | 2   |
| Total                      | 2912*| 2314*| 318| 124| 117 | 16  | 12  | 10  |

* For French samples, one information is missing (adults/veal), one is added to total.

### Table 2. The overall seroprevalence of *Toxoplasma gondii* infection in beef consumed in France accordingly to the terminal titer (6; 10; 25; 50; 100; 200). Only one positive sample at terminal titer of 400 (adult of French origin).

| All Mean and 95% CI | 6 | 10 | 25 | 50 | 100 | 200 |
|---------------------|---|----|----|----|-----|-----|
| Samples of French origin (N = 2349*) | 16.96 [11.91–23.56] | 7.77 [4.91–12.09] | 3.93 [1.90–7.95] | 1.58 [0.54–4.52] | 0.96 [0.30–3.05] | 0.52 [0.12–2.17] |
| Imported samples (N = 563) | 19.75 [7.88–41.48] | 15.52 [6.32–33.32] | 11.15 [3.54–30.04] | 0.57 [0.07–4.33] | 0.38 [0.05–2.91] | 0.38 [0.05–2.91] |
| Overall (N = 2912*) | 17.38 [12.73–23.26] | 8.93 [5.97–13.17] | 5.01 [2.73–9.02] | 1.43 [0.53–3.81] | 0.87 [0.30–2.49] | 0.50 [0.14–1.74] |

Note. % prevalence, 95% CI: 95% confidence interval of prevalence.

* One information missing (adults/veal), one is added to total.
Starting from the top:

### Table 3

The overall seroprevalence of *Toxoplasma gondii* infection in calves consumed in France accordingly to the terminal titer (6; 10; 25; 50; 100; 200).

| Area of slaughtering | All Mean and 95% CI | 6 | 10 | 25 | 50 | 100 | 200 |
|----------------------|---------------------|---|----|----|----|-----|-----|
| Samples of French origin (N = 574) | 5.06 [1.79–13.50] | 2.44 [0.35–14.94] | 1.27 [0.17–8.89] | 0.61 [0.08–4.68] | 0.01 [0.00–0.12] | 0 |
| Imported samples (N = 225) | 6.34 [4.94–8.10] | 5.17 [3.28–8.06] | 3.67 [1.59–8.22] | 0 | 0 | 0 |
| Overall (N = 799) | 5.34 [2.49–11.09] | 3.03 [0.89–9.82] | 1.78 [0.56–5.51] | 0.48 [0.06–3.55] | 0.01 [0.00–0.09] | 0 |

### Table 4

The overall seroprevalence of *Toxoplasma gondii* infection in adults consumed in France accordingly to the terminal titer (6; 10; 25; 50; 100; 200).

| Area of slaughtering | All Mean and 95% CI | 6 | 10 | 25 | 50 | 100 | 200 |
|----------------------|---------------------|---|----|----|----|-----|-----|
| Samples of French origin (N = 1774) | 22.00 [16.16–29.22] | 10.03 [7.22–13.79] | 5.05 [2.76–9.07] | 1.99 [0.76–5.10] | 1.36 [0.42–4.34] | 0.74 [0.17–3.11] |
| Imported samples (N = 338) | 31.41 [19.23–46.83] | 24.51 [15.38–36.72] | 17.65 [7.51–36.14] | 1.07 [0.20–5.51] | 0.71 [0.13–3.71] | 0.71 [0.13–3.71]* |
| Overall (N = 2112) | 23.12 [17.83–29.40] | 11.75 [8.51–16.00] | 6.55 [3.78–11.12] | 1.88 [0.78–4.46] | 1.28 [0.45–3.64] | 0.73 [0.21–2.54] |

* No terminal titer at dilution 100 (but at 200 or 50).

### Table 5

Seroprevalence of *Toxoplasma gondii* infection (cut-off titer 6) in bovine meat consumed in France by area of slaughtering.

| Area of slaughtering | Bovine (adults and calves) | % (No. of animals sampled) | 95% CI | Adults | % (No. of animals sampled) | 95% CI | Calves | % (No. of animals sampled) | 95% CI |
|----------------------|-----------------------------|-----------------------------|-------|--------|-----------------------------|-------|--------|-----------------------------|-------|
| Alsace-Franche Comté | 9.92 (199) | 3.01–28.08 | 33.53 (70) | 14.88–59.29 | 5.61 (129) | 2.01–14.68 |
| Aquitaine Midi-Pyrénées | 16.02 (150) | 13.14–19.38 | 25.28 (76) | 15.16–39.04 | 7.27 (74) | 4.08–12.63 |
| Auvergne-Limousin | 3.64 (150) | 0.51–21.70 | 15.60 (67) | 7.77–28.87 | 0 (83) | 0 |
| Bourgogne | 12.59 (100) | 8.05–19.16 | 13.39 (93) | 9.52–18.51 | 0 (7) | 0 |
| Bretagne | 12.01 (400) | 6.91–20.07 | 16.84 (269) | 10.67–25.57 | 0 (131) | 0 |
| Centre-Poitou Charentes | 30.79 (100) | 11.58–60.20 | 29.65 (91) | 11.51–57.71 | 55.56 (9) | 21.12–85.37 |
| IDF-Picardie-Champagne-Ardennes | 5.61 (150) | 2.86–10.71 | 5.95 (139) | 3.10–11.15 | 1.49 (11) | 0.11–17.10 |
| Languedoc-Roussillon-PACA | 23.33 (100) | 17.35–30.61 | 27.93 (80) | 17.22–41.93 | 15.00 (20) | 4.92–37.59 |
| Lorraine | 22.49 (150) | 8.17–48.60 | 23.66 (143) | 8.89–49.62 | 0 (7) | 0 |
| Nord-Pas-de-Calais | 27.48 (100) | 19.17–37.72 | 30.98 (90) | 23.82–39.19 | 0 (10) | 0 |
| Normandie | 26.16 (200) | 9.29–59.31 | 24.25 (167) | 8.55–52.29 | 34.02 (33) | 6.27–79.88 |
| Pays de la Loire | 21.19 (400) | 9.65–40.35 | 22.66 (377) | 10.39–42.53 | 0 (23) | 0 |
| Rhône-Alpes* | 28.73 (150*) | 8.10–64.82 | 35.31 (112) | 9.99–72.86 | 5.94 (37) | 1.48–20.97 |
| Bovine meat consumed in France (French slaughterhouses)* | 16.96 (2349*) | 11.91–23.56 | 22.00 (1774) | 16.16–29.22 | 5.06 (574) | 1.79–13.50 |

Note. % prevalence, 95% CI: 95% confidence interval of prevalence.
* One information missing (adults/veal), one is added to total.

Cut-off was chosen. The effect of age as a qualitative variable (calves/adults) in univariate analysis was found to be highly significant (p = 0.00081).

Depending on the region of slaughter, seroprevalence varied between 0% and 55.56% for calves and from 5.95% to 35.31% for adults (Table 5), with no specific geographical pattern identified (Fig. 2A–C). The seroprevalence estimates were markedly lower in calves than in adults, in all regions, except Centre, Poitou-Charentes and Normandie. Seroprevalence estimates by areas of birth are also given in Table 6, with the same identified difference between calves and adults, which stands for all regions, except IDF, Picardie, and Champagne-Ardennes.

The terminal titer of the MAT, in relation to age is presented in Figure 3 for all French samples (n = 2348) (age in years) (A) or only for bovines less than 1 year old (n = 601) (age in months) (B). No particular trend was observed, with higher or lower terminal titer specifically for young or old animals. The seroprevalence of *T. gondii* infection in bovines of French origin (adults + calves) according to the terminal titer (6; 10; 25; 50; 100; 200) is presented in Figure 4. The seroprevalence showed an increase in general at a younger age;
however, after the age of 4–5 years, a general linear trend was not clearly observed.

**Imported samples**

The overall seroprevalence in imported beef at a cut-off titer of 6 was estimated to be 19.75% [7.88–41.48%] (Tables 2 and 7), with a distinct seroprevalence of 6.34% [4.94–8.10%] for calves (Tables 3 and 7) and 31.41% [19.23–46.83%] for adults (Tables 4 and 7). As found previously, the seroprevalence decreased for both categories, when a higher cut-off was chosen (Tables 2–4). The higher seroprevalence in imported animals compared to French origin bovines, for any of the chosen titers (Tables 2–4), was not found to be significant in univariate analysis, whether for bovines overall (adults + calves) \( (p = 0.70) \), or for adults \( (p = 0.13) \) or calves \( (0.65) \), respectively.

**Risk factor analysis in samples of French origin**

The analysis was restricted to animals below 12 years old since the number of observations above or equal to 12 years was 3% in the dataset, and in order to avoid fitting a nonlinear relationship with a small number of samples. The complete dataset for all explanatory variables, such as age, sex, type of production (milk or meat), area of birth, breeding and slaughtering, contained 1642 observations.

In order to investigate different risk factors, 195 possible models were tested. The best model with the lowest Akaike’s Information Criterion \( (AIC = 1453.2) \) included only age as an explanatory variable. This final model is a polynomial model of degree 4 with age in years \( (p = 0.70) \), showing also that a linear model was not the best model. It is also the most parsimonious model for this range of AIC. Other models with differences in AIC lower than 2, included a polynomial of degree 4 for age in years and sex in interaction \( (AIC = 1453.66) \), and a polynomial of degree 5 for age in years \( (AIC = 1453.8) \). Figure 5 shows the relative agreement between observed values for age in class by year and the prediction of the final model for the mean age for each annual class. Except for the age class 6 (5.9 years in average) and 7 (7.1 years in average) years old, the final model fits the data: the seroprevalence for each particular age class, represented by the top of the grey bar, is inside the 95% confidence interval of the prediction of the model. Figure 6 shows the prediction of the final model, with a clear non-linear effect of age related seroprevalence (at a cut-off titer of 6).

**Toxoplasma gondii isolation and genotyping**

The digestion products of 209 heart samples, obtained from both serologically positive \( (n = 200) \) and negative \( (n = 9) \) animals, were inoculated in mice. In total, 627 mice were used for the bioassay. For four bovine samples (8, 76, 122, 125 months; terminal titer of 10, 10, 25 and 10, respectively), all three mice died in the first 48 h post-inoculation (p.i.), yielding 205 bioassay results. For two samples, corresponding to serologically

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Figure 2. Geographical variation of *Toxoplasma gondii* seroprevalence of French bovine samples according to the area of slaughtering and to age categories: (A) calves; (B) adults; (C) bovines overall (calves and adults). The numbers represent the number of samples collected for each region.
positive adult bovines (4.8 and 7 years old; terminal titer of 25 and 10), collected in Franche-Comté and Languedoc-Roussillon region slaughterhouses, we noted seroconversion in the inoculated mice 4 weeks p.i.; a microscope examination at 60 days p.i. confirmed the presence of tissue cysts in brains. Both strains were genotyped as type II with all 15 microsatellite markers (TUB2, W35, TgM-A, B18, B17, M33, MIV.1, MXI.1, M48, M102, N60, N82, AA, N61, N83) being

Table 6. Seroprevalence of *Toxoplasma gondii* infection (cut-off titer 6) in bovine meat consumed in France by area of birth.

| Area of slaughtering | Bovine (adults and calves) | Adults | Calves |
|----------------------|-----------------------------|--------|--------|
|                      | % (No. of animals sampled)  | 95% CI | % (No. of animals sampled) | 95% CI | % (No. of animals sampled) | 95% CI |
| Alsace-Franche Comté | 12.38 (60)                  | 7.04–20.86 | 14.25 (36) | 8.62–22.65 | 7.81 (24) | 1.45–32.85 |
| Aquitaine Midi-Pyrénées | 13.22 (121)            | 5.73–27.64 | 20.45 (63) | 10.64–35.70 | 4.32 (58) | 0.79–20.38 |
| Auvergne-Limousin    | 16.38 (197)                 | 8.34–29.67 | 30.55 (113) | 17.45–47.79 | 2.96 (84) | 0.94–8.91 |
| Bourgogne            | 30.95 (111)                 | 15.09–53.06 | 34.35 (99) | 16.51–58.08 | 3.57 (12) | 0.55–19.95 |
| Bretagne             | 17.96 (206)                 | 7.80–36.16 | 24.43 (126) | 11.48–44.61 | 9.87 (80) | 1.54–43.40 |
| Centre-Poitou Charentes | 14.02 (228)            | 6.97–26.20 | 24.51 (144) | 13.99–39.34 | 4.51 (84) | 1.34–14.11 |
| IDF-Picardie-Champagne-Ardennes | 5.40 (141) | 2.01–13.71 | 5.15 (122) | 1.54–15.86 | 6.47 (19) | 1.44–24.65 |
| Languedoc-Roussillon-PACA | 17.85 (64)           | 11.23–27.19 | 30.78 (54) | 15.25–52.35 | 3.74 (10) | 0.40–27.17 |
| Lorraine             | 3.84 (101)                  | 0.62–20.36 | 4.25 (97)  | 0.67–22.63 | 0 (4) | 0 |
| Nord-Pas-de-Calais   | 23.80 (50)                  | 17.62–31.32 | 29.96 (39) | 25.44–34.90 | 1.05 (11) | 0.11–9.10 |
| Normandie            | 15.07 (249)                 | 8.09–26.35 | 15.84 (228) | 8.79–26.87 | 6.01 (21) | 0.77–34.42 |
| Pays de la Loire     | 20.07 (349)                 | 10.95–33.91 | 25.24 (278) | 14.28–40.62 | 4.26 (71) | 0.95–17.14 |
| Rhône-Alpes*         | 20.44 (113)                 | 9.28–39.22 | 25.60 (74)  | 10.49–50.25 | 6.59 (39) | 1.26–28.07 |
| Bovine meat consumed in France (French slaughterhouses)* | 16.96 (2349*) | 11.91–23.56 | 22.00 (1774) | 16.16–29.22 | 5.06 (574) | 1.79–13.50 |

Note. % prevalence, 95% CI: 95% confidence interval of prevalence.
* One information missing (adults/veal), one is added to total.

Figure 3. Terminal titer of the modified agglutination test (MAT) for French origin samples in relation to age (A) for all samples (*n* = 2348) (age in years); (B) only for bovines less than 1 year (*n* = 601) (age in months). The number of observations at each month of age is given at the top of the corresponding bar.
amplified. They can be found in the *Toxoplasma* Biological Resource Center, Reims, France (toxocrb.com) under accession numbers: TgA 32127 and TgA 32128. No strains were isolated from the heart samples corresponding to the nine serologically negative bovines. Detailed data about the number of collected and bioassayed samples of French origin according to the terminal titer + and age can be found in the Supplementary file.

### Discussion

Following a previous study and using a sampling plan based on ovine meat production data and innovative testing for antibody detection and parasite isolation [33], the present survey, conducted according to the same principles of sampling and testing, aimed at investigating the national *T. gondii* seroprevalence in beef consumed in France. Therefore, the

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**Figure 4.** Seroprevalence of *Toxoplasma gondii* infection in bovines of French origin (adults + calves) accordingly to the age and the titer (6; 10; 25; 50; 100; 200).

**Table 7.** Seroprevalence of *Toxoplasma gondii* infection (cut-off titer 6) in bovine meat imported at IFM Rungis (Ile de France).

| Origin country          | Relative frequency in French meat imports | Bovine (adults and calves) | Adults | Calves |
|-------------------------|-------------------------------------------|-----------------------------|--------|--------|
|                         |                                           | % (No. of animals sampled) | 95% CI | % (No. of animals sampled) | 95% CI | % (No. of animals sampled) | 95% CI |
| The Netherlands          | 25.1%                                     | 5.88 (119)                  | 2.83–11.83 | No data (0) | No data | 5.88 (119) | 2.83–11.83 |
| Germany                  | 24.7%                                     | 38.46 (143)                 | 30.85–46.68 | 40.91 (132) | 32.85–49.49 | 9.09 (11) | 1.26–43.90 |
| Ireland                  | 15.3%                                     | 23.81 (84)                  | 15.90–34.06 | 23.81 (84) | 15.90–34.06 | No data (0) | No data |
| Italy                    | 10.1%                                     | 5.63 (71)                   | 2.13–14.08 | 0 (1) | 0 | 5.71 (70) | 2.16–14.27 |
| Belgium                  | 8.4%                                      | 18.46 (65)                  | 10.79–29.77 | 20.00 (60) | 11.72–32.01 | 0 (5) | 0 |
| Poland and other European countries* | 4.8%                                      | 9.52 (21)                   | 2.39–31.15 | 0 (1) | 0 | 10.00 (20) | 2.51–32.41 |
| UK                       | 1.5%                                      | 30.30 (33)                  | 17.14–47.76 | 30.30 (33) | 17.14–47.76 | No data (0) | No data |
| Austria                  | 0.8%                                      | 33.33 (27)                  | 18.33–52.69 | 33.33 (27) | 18.33–52.69 | No data (0) | No data |
| Total imports            | 90.7%                                     | 19.75 (563)                 | 7.88–41.48 | 31.41 (338) | 19.23–46.83 | 6.34 (225) | 4.94–8.10 |

* Poland (13) + Denmark (1) + Lithuania (4) + Latvia (1) + Czech Republic (1) + Slovakia (1).
calculated seroprevalence should be considered the most representative value for *T. gondii* in beef in France, originating either from slaughtered or imported carcasses. Currently, France is one of the largest producers of beef in Europe, but the import of 300,000 tons per year remains necessary [12]. Compared to what was initially planned for the collected samples from imports, several countries were absent, in particular Spain and Brazil, representing 7.5% and 0.8% of the relative frequency in French meat imports. This explains the 90.7% of the total imports, which represents an acceptable level of sample collection.

The overall seroprevalence of *T. gondii* in bovine carcasses was 17.38% [12.73–23.26] at a cut-off titer of 6. This is in accordance with previous results obtained in France [31, 52], in other European countries (Spain: 18.6% [4], Estonia 18.62% [38]), or elsewhere in the world (Senegal: 13% [17]), with the same technique and testing a high number of samples (>1000). However, seroprevalence in cattle varies greatly throughout the world, ranging from 0% to 92% [2, 10, 21, 26, 31, 36–41, 45, 47, 50, 51, 55, 57]. This variation is due to a large number of factors, such as climate, herd-management, cat population, sampling procedure, or diagnosis tools [31, 40, 47]. Therefore, in order to introduce less bias for comparison between studies, we used the MAT technique, which has already been demonstrated to be superior to other tests (Sabin-Feldman Dye Test, latex agglutination, indirect haemagglutination) for serological screening of cattle [22]. The choice of a low threshold (cut-off titer of 6) was justified in our case by (a) our goal: to isolate viable *T. gondii* parasites, considering at that time that viable parasites should mostly be present in seropositive animals; (b) the fact that generally in bovines one can find very low titers, in our case the highest titer was observed in only one bovine at a terminal titer of 400, most of the animals (559/598) having a titer lower than 50; and (c) the decision to work on muscle (cardiac/diaphragm) fluid, representing a low-concentration antibody matrix compared to serum. Therefore our titer corresponds to a higher titer in blood sera. A 10-fold equivalence between the two matrices has been suggested in a study comparing meat juice and serum for *Trichinella* detection [46].

### Table 8. Final model to explain *Toxoplasma gondii* seropositivity at a cut-off titer of 6 (1642 observations).

| Factor | Parameter estimate (Standard Error) | OR   | 95% CI OR | P value (Wald Test) |
|--------|------------------------------------|------|-----------|---------------------|
| Intercept | −3.547 (0.657) | 0.029 | 0.008–0.104 | 2.37 × 10⁻⁵ |
| Age²   | 2.190 (0.694) | 8.931 | 2.293–34.783 | 0.0047 |
| Age³   | −0.659 (0.254) | 0.517 | 0.315–0.851 | 0.0169 |
| Age⁴   | 0.076 (0.034) | 1.079 | 1.010–1.153 | 0.0340 |
| Age⁵   | −0.003 (0.001) | 0.997 | 0.994–1 | 0.0551 |

**Figure 5.** Comparison of observed values versus predicted values by the final model according to age. The observed values are in green bars, while for the predicted values the red point represents the mean prediction and the blue segment the 95% confidence interval of the prediction. The number above the blue segment is the number of observations for this particular class of age.
A strong age (adult/calves) effect was observed with a seroprevalence of 5.34% [2.49–11.09] for calves (animals less than 8 months of age) and 23.12% [17.83–29.40] for adults (animals over 8 months of age) at a cut-off titer of 6. The age difference was valid for both types of samples: French and imported, and for all French regions except Normandie and Centre-Poitou Charentes, where a higher seroprevalence was observed in calves than in adults. However, regarding confidence intervals this difference was not meaningful. This difference could be speculated to be due to the low number of animals (33 and 9, respectively) or to a common source of infection. Differences in seroprevalence between calves and adults may be related to the oral route of infection, older animals being exposed to the parasite for a longer period of time than younger ones. The overall seroprevalence and the significant age difference in the seroprevalence in cattle, follows the same pattern that has already been observed in France for other meat-producing animals [19, 33]. Nevertheless, the percentages were significantly higher for other ruminants (sheep), both for adult meat (81%) and lamb (15%).

The collection of French samples, and their associated data, enabled us to analyse the effect of several parameters: age and sex of animals, geographical origin and bovine meat production level in the area where the sample was collected. Other parameters such as seasonal variations, herd characteristics and herd management were not taken into account because of several reasons: the timeframe was limited (6-month period); the goal of our study was to estimate seroprevalence of *T. gondii* infection in fresh beef consumed in France and not in the cattle population of France. Nevertheless, statistical analyses reveal that the major factor to explain variations in the seroprevalence observed is the age of the animal at a threshold dilution of 1:6 with a non-linear relationship (polynomial of degree 4). A non-linear relationship was also found graphically between age and seroprevalence at any cut-off (Figs. 4–6), suggesting no life-long immunity and possible re-infection by *T. gondii* in cattle. Similar age-related seroprevalence patterns were obtained in a recent study in Estonia [38], where the seroprevalence increased until the age of 5 years (60–71 months), after which there was no evident linear pattern. One possible explanation might be that antibodies do not persist in cattle, already shown by Dubey and Thuliez [26] and Gilot-Fromont et al. [31], while antibodies could persist several years in sheep [27], pigs [19], or other animals [28]. Another explanation might be that seropositive animals were culled earlier, therefore *T. gondii* infection might be negatively associated with longevity [38]. Moreover, it was shown that the presence of cats modified the age-seroprevalence relationship [31]. This factor could not be tested here, and our population probably mixed animals with life-history presence of cats and absence of cats. At the same time, no clear relationship was detected between the terminal titer and age, with calves having low titers, adults having high titers, or the other way around (Fig. 3A and B). In our population, the high titers (>50) were found in animals less than 10 years old, together also with low titers, while in older animals (>10 years), we only found low titers (<50). To our knowledge, no other recent study has investigated this pattern, so only speculations about the loss of immunity in older cattle might explain this, as previously mentioned [26, 31]. Concerning the animals younger than 1 year, one can clearly see that calves around 3–4 months are positive at a low titer (6). However, this result was for a small number of animals, while animals between 6 and 9 months were found to be positive at various titers (high and low), suggesting the detection/presence of maternal antibodies (3–4 months) and the first *T. gondii* infection (6–9 months), with the decrease of maternal immunity. Similar situations with the decrease of maternal antibodies and the first natural *T. gondii* infection revealed by higher antibodies titers are found in other animal species, such as sheep [60] or pigs [19]. When comparing the seroprevalence estimates given by area of slaughtering (Table 5) and area of birth (Table 6), a variation was observed, showing that the areas with the highest seroprevalence were not the same between these two variables. This is true for adults and bovines overall (calves + adults), while for calves the number of positive analyses was too low to continue the comparison. One of the hypotheses to explain this variation is the commercial trade: a similar seroprevalence between birth and slaughtered area should be found only at a local level, not in the case of a national survey, when samples are collected from animals that are potentially born in region A, raised in region B, and slaughtered in region C. This hypothesis is also supported by a recent UK study [37], showing that the number of sites an animal stayed before slaughter ranged from 1 to 15 (median 3), thereby multiplying the possibilities of *T. gondii* oocyst ingestion, and making it relatively impossible to identify the infection site. A similar situation might also be present in France. Moreover, the age of slaughtering can be seen as a confounding factor for comparing different areas.

Our study enabled us to observe that several regions (Aquitaine, Midi Pyrénées, Nord-Pas de Calais), for which the seroprevalence was high in adult animals, and that provide meat that is eaten raw or undercooked (*bleu* or *saignant*) (Fig. 2B), corresponded to those for which seroprevalence was also high in humans [10, 58]. Our results are in agreement with the previous study of Halos et al. [33], that found similar areas of concordance/correlation between seroprevalence of *T. gondii* in humans and lamb, the lamb representing the other
type of meat that is often eaten undercooked (agneau rosé) in France. No direct link can yet be made between the presence of *T. gondii* in meat-producing animals and human infection, except if we assume that locally produced meat is consumed locally (very unlikely). On the other hand, other regions with high seroprevalence in adult animals (Rhone-Alpes, Centre-Poitou Charentes) do not present a high seroprevalence in humans. Therefore, the agreement between *T. gondii* seroprevalence in humans and cattle in several areas of France should be interpreted carefully, as it remains unexplained. It could be speculated to be due to more intense circulation of the parasite in certain areas, but this requires further investigation.

Concerning the bioassays and parasite isolation, several parameters need to be taken into account: first, only a limited number of samples (209/2349) could be tested, thus reducing our analysis on the prevalence of viable *T. gondii* tissue cysts in beef consumed in France. Second, bioassays in mice are less sensitive than those in cats [24], especially in cases where the whole heart digest could not be fully inoculated. Third, our results probably underestimate the effective parasitological prevalence in beef, since we mostly tested serologically positive animals (200/209). This was done based at that time on the results from the ovine survey [33], where the chance of obtaining viable parasites increased with the titer and age. Meanwhile, in a study on a limited number of bovines (n = 100), the authors identified two PCR-positive results, while the corresponding MAT results were negative (cut-off 1:40). Similarly, three MAT-positive results had corresponding negative PCR results [49]. In another recent study on 167 calves and 235 adults, the authors demonstrated that serological testing by MAT or p30 immunoblot does not provide reliable information about the presence of *T. gondii* parasites or DNA in cattle [50]. In our case, however, two strains (1%) were isolated from heart samples of low titer MAT-positive cattle, adding to the limited number of such reports [7, 25, 50, 51]. The presence of live parasites in the heart of naturally infected cattle does not indicate their presence in other parts of carcasses that will be consumed. Nevertheless, previous studies showed persistence of viable *T. gondii* in tissues likely to be eaten, including the heart, after experimental infection with oocysts [26]. Accurate knowledge of the distribution of the parasite within cattle carcasses is still needed [16, 18, 28, 29]. Similar results have previously been obtained for the ovine meat consumed in France, with a significantly higher prevalence of parasite isolation (5.4%) from fresh carcasses [33].

The animals in this study were sampled in 2009, and it should be emphasized that the epidemiological situation may have changed since then. Nevertheless, the samples were collected and analysed in a comparative way with the national surveys of sheep and pigs [19, 33]. Therefore, the present study adds to knowledge on the presence of *T. gondii* in the three main meat producing animals in France, offering new and reliable facts about *T. gondii* presence in bovines. Overall, our results indicate that there is a risk of human infection with *T. gondii* following beef consumption in France, which still needs to be correlated with the food habit of ingesting raw/undercook (bleu or saignant) beef [9, 58]. However, new questions have emerged specifically concerning the isolation of parasites from beef and the precise role of bovines, generally described as poor hosts for *T. gondii*, in human infection. Similar surveys should be conducted in other countries, to obtain a broader view of animal sources of *T. gondii* infection in humans.

### Supplementary materials

**Table.** Number of collected/bioassayed samples of French origin, according to the terminal titer and the age of animals.

[Supplementary material available at https://www.parasite-journal.org/10.1051/parasite/20190766olm](https://www.parasite-journal.org/10.1051/parasite/20190766olm)

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### Conflict of interest

All individual authors declare that they have no conflict of interest (financial, personal or other).

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