





Caratterizzazione genomica di Campylobacter e sviluppo di un modello per la tracciabilità del microrganismo da diverse fonti

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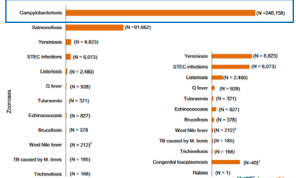
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# Campylobacter in EU







doi: 10.2903/j.efsa.2018.5500

### The European Union summary report on of zoonoses, zoonotic agents and food-t 2017

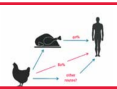
European Food Safety Authority and European Centre for Disa (EFSA and ECDC)

Table 2: Reported hospitalisation and case fatalities due to zoonoses in confirmed human cases in the EU, 2017

Disease	Number of confirmed <sup>(a)</sup> Human cases	Hospitalisation				Deaths			
		Status available (%)	Number of reporting MS <sup>(b)</sup>	Reported hospitalised cases	Proportion hospitalised (%)	Outcome available (%)	Number of reporting MS <sup>(b)</sup>	Reported Deaths	Case Fatality (%)
Campylobacteriosis	246,158	27.6	17	20,810	30.5	72.8	16	45	0.04
Salmonellosis	91,662	43.1	14	16,796	42.5	67.8	17	156	0.25
Yersiniosis	6,823	27.1	14	616	33.4	65.5	15	3	0.07
STEC infections	6,073	41.0	18	933	37.5	66.1	21	20	0.50
Listeriosis	2,480	40.4	16	988	98.6	65.8	18	225	13.8
Q-fever	928	NA <sup>(c)</sup>	NA	NA	NA	56.0	10	7	1.35
Echinococcosis	827	31.2	14	140	54.3	30.1	14	1	0.40
Brucellosis	378	45.8	10	104	60.1	33.9	10	1	0.78
Tularaemia	321	38.3	9	76	61.8	51.1	9	1	0.6
West Nile fever(a)	212	72.2	8	134	87.6	98.6	9	25	12.0
Trichinellosis	168	44.6	9	56	74.7	40.5	9	0	0.0
Congenital toxoplasmosis	40	57.9	3	18	NA	63.2	3	0	0.0
Rabies	1	NA(c)	NA	NA	NA	0.0	0	NA	NA

# Inferring host-host transmission: zoonotic transmission of *Campylobacter jejuni*









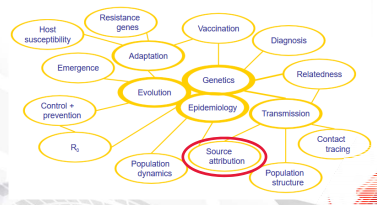






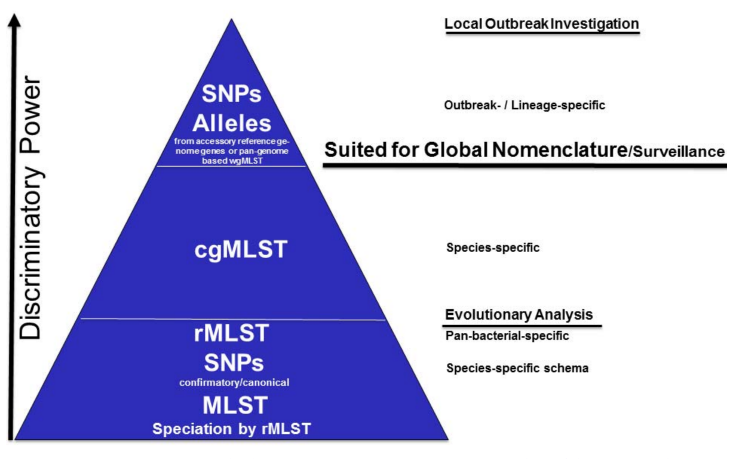


# Evolutionary genetics as a framework for understanding genetic diversity



- Per stimare i contributi relativi di fonti differenti (animali, ambientali o alimentari) alla malattia nell'uomo;
- -guidare degli interventi mirati e misurera loro impatto;
- -diversi approcci, tra cui

MICROBIAL SUBTYPING

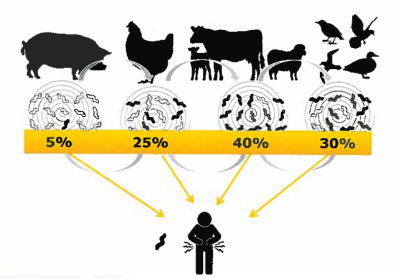


Bassa sensibilità Molto robusta Buona per analisi filogenetiche Non abbastanza per investigare focolai

# Analisi dell'attribuzione alle fonti



- Asymmetric Island Model (Wilson et al. PLoS Genet 2008;4:e1000203)
  - Bayesian population genetics model that uses MLST data for probabilistic assignment of STs to sources
  - Campylobacter population is seen as separate islands where mutation, recombination and migration occur



### **Evolutionary stage**

- · Only source data
- Estimates mutation, recombination and migration rates to be used in the second stage

### Attribution stage

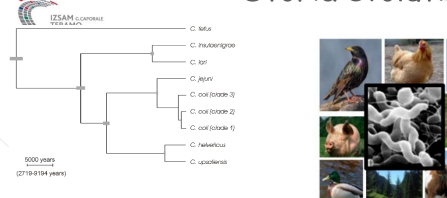
- Relies on sampling probabilities: probability that a given human ST is randomly sampled from each host population
- Estimates an assignment posterior probability for each ST

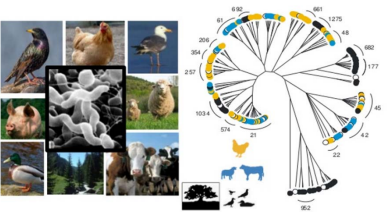
Lapo Mughini Gras

Infezioni da Campuloharter pell'uomo pai Rosci Rosci Lya Dicambra accan

# Storia evolutiva

# C.jejuni



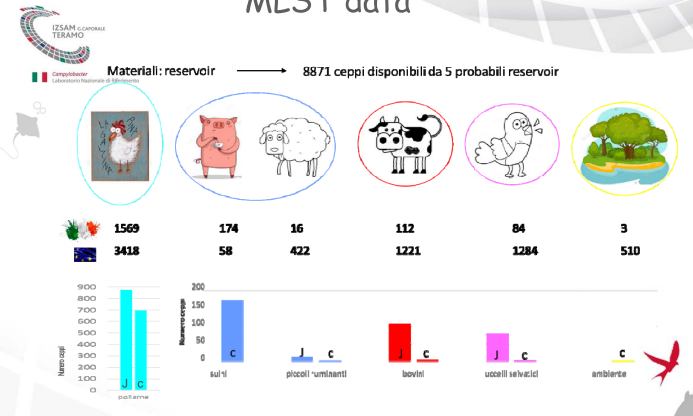


C. jejuni si caratterizza per la sua diversità genomica e per possedere una struttura di popolazione non clonale dovuta alla presenza di elevata ricombinazione genetica

Sheppard et al. (2009) Clinical Infectious Diseases 48:1072–1078 Sheppard et al. (2010) Applied Environmental Microbiology 76, 5269-5277

# Clade 2 Sheppard SK, Dallas JF, Wilson DJ, Strachan NJC, McCarthy ND, et al. (2010) Evolution of an Agriculture-Associated Disease Causing Campylobacter coli Clade: Evidence from National Surveillance Data in Scotland, PLOS ONE 5 (12): e15708. https://doi.org/10.1371/journals.plos.org/plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plosone/article?id=10.1371/journals.plos

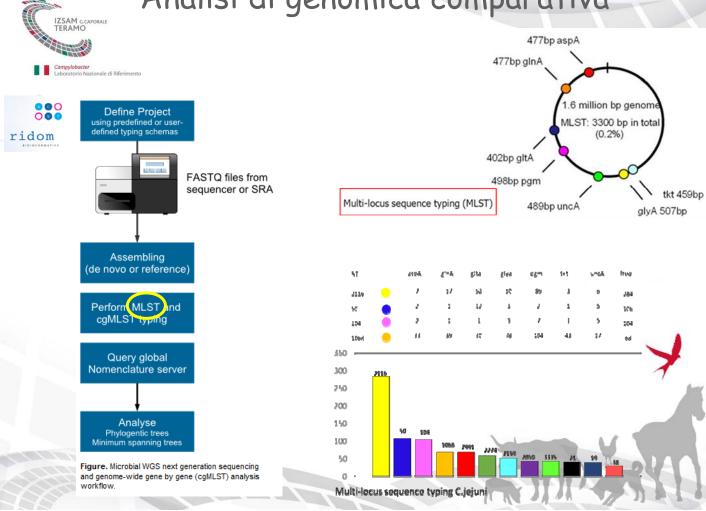
# MLST data

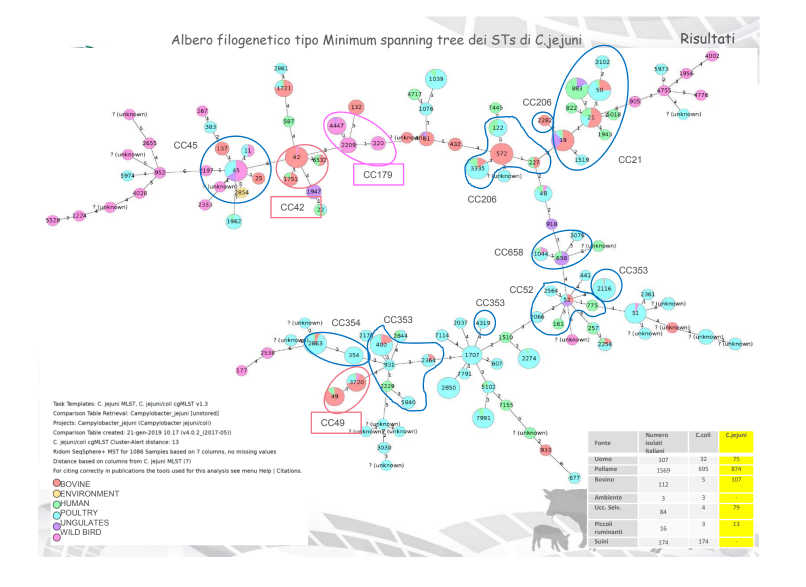


107 humans (75 c.jejuni-32 C.coli)

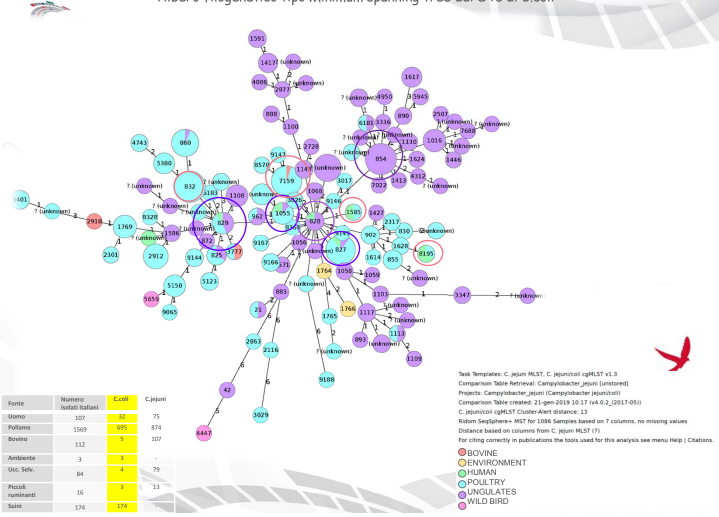


# Analisi di genomica comparativa



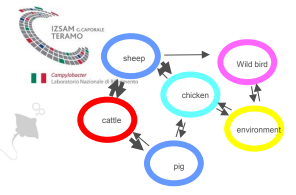


### Albero filogenetico tipo Minimum spanning tree dei STs di C.coli



## Source attribution

### Risultati

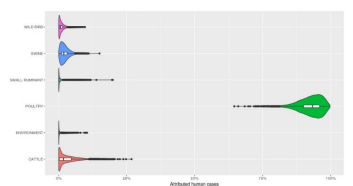


frontiers
in Microbiology

ORIGINAL RESEARCH published: 13 June 2016 10 3389/fmich 2016 00887

# Tracing Back Clinical Campylobacter jejuni in the Northwest of Italy and Assessing Their Potential Source

Elisabetta Di Giannatale¹, Giuliano Garofolo¹, Alessandra Alessiani¹, Guido Di Donato¹, Luca Candeloro², Walter Vencia², Lucia Decastelli³ and Francesca Marotta¹\*



### https://github.com/jmarshallnz/islandR

Wilson et al 2008

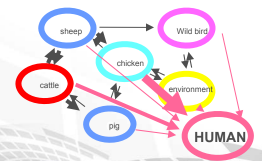


### islandR

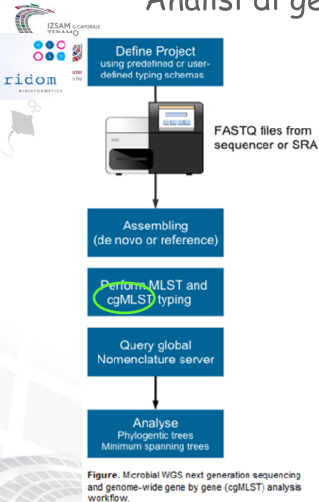
The islandR package allows source attribution using the island genomic model (and in the future, other genomic models).

This allows the attribution of cases of disease to their likely sources.

The assymmetric island model estimates the sampling distribution of genotypes on sources by using the genetic distance between isolates to infer mutation and recombination rates in addition to migration rates between sources. This allows improved estimation of the sampling distribution over and above that achieved using just the prevalence of each type. Further, it allows estimating the likely prevalence of unobserved genotypes.

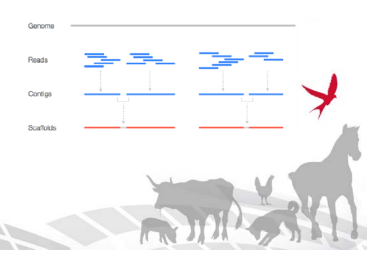


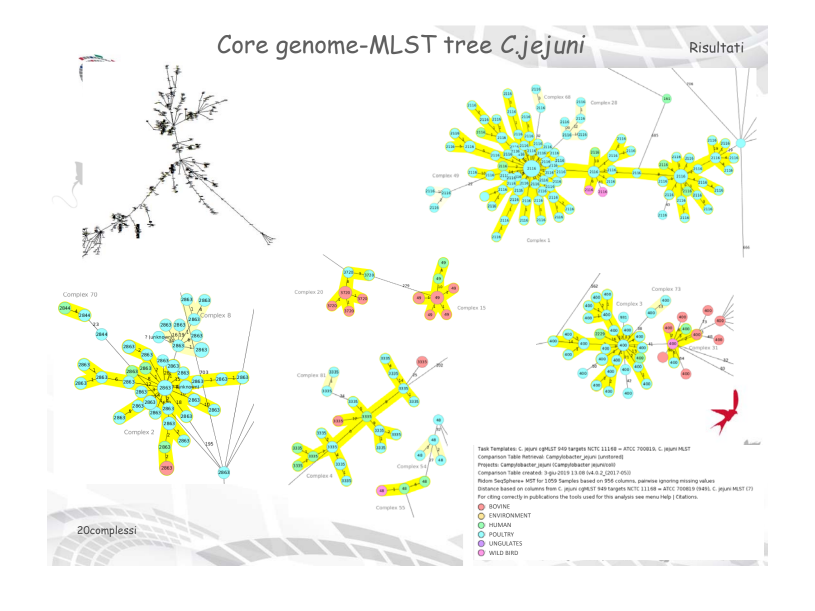
# Analisi di genomica comparativa

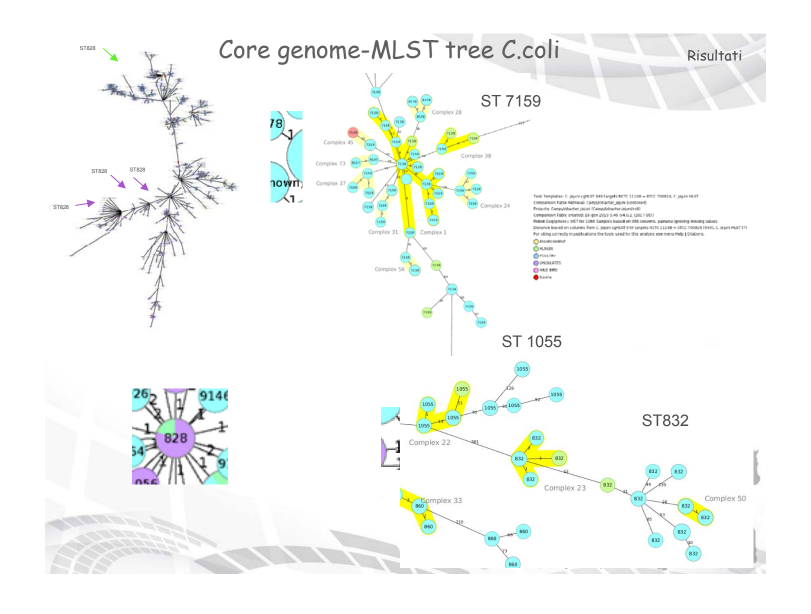




- Library prep (Nextera XT DNA Library Preparation Kit-Illumina)
- Sequencing (NextSeq500 –Illumina)
- De novo assembly (SPAdes)



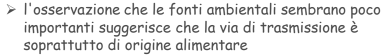












- > interventi mirati sulla produzione primaria dei polli al fine di ridurre l'incidenza della malattia.
- grande variabilità esistente anche nei ceppi di Campylobacter isolati dagli umani, molteplici foci d'infezione.



WGS è un potente strumento per la sorveglianza di agenti patogeni di origine alimentare, stabilire un legame tra casi non correlati basati solo su livelli di prossimità della distanza genomica potrebbe essere fuorviante.







### Elisabetta Di Giannatale

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