Comparative genomics provides insights into the evolution of novel *Fusarium oxysporum* f. sp. *lactucae* race variants

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1. Introduction

- Fusarium oxysporum f. sp. lactucae (Fola) is a devastating soil-borne pathogen that causes Fusarium wilt of lettuce, leading to crop losses exceeding 50% worldwide.
- The recent emergence and spread of the highly aggressive Race 4 across Northern Europe poses a significant new threat to both protected and open field lettuce production.
- Initial molecular and pathological characterisation of field isolates revealed unique variants (v), including a potential new race (NR) from France capable of breaking multiple sources of genetic resistance.



2. Methods

- We performed Nanopore long-read sequencing to generate high quality genome assemblies for 16 diverse Fola isolates, including variants (v) of Race 1 (R1) and 4 (R4), a potential new race (NR), and the first assemblies for Race 2 (R2) and 3 (R3) (Figure 1).
- Phylogenomic analysis of 3,804 single-copy orthologous genes was conducted to resolve the evolutionary relationships between Fola races.
- Comparative genomics and the FoEC2 pipeline were used to analyse the diversity of accessory regions and identify the repertoire of putative effector genes across all isolates.



3. Results

- Phylogenomic analysis reveals that Fola is **polyphyletic.** R1 and R4 cluster closely, but R2, R3, and the potential new race (NR) are distributed separately throughout the *F*. oxysporum phylogeny (Figure 2).
- The potential new race (NR) isolate possesses an accessory region that is distantly related to the four previously described races and can overcome both R1 and R4 resistance in lettuce differentials (Figure 2).
- We identified a core set of 44 effectors common to all isolates (PanFola), but also racespecific effectors, with the new race (NR) isolate containing 4 unique effectors not shared with any other Fola isolates (Figure 3).



4. Discussion

- The polyphyletic ancestry of Fola indicates that the ability to cause disease on lettuce has evolved independently multiple times within the *F. oxysporum* species complex.
- Distinct profiles of putative effector genes, especially those unique to specific isolates, likely underpin the observed differences in pathogenicity and the ability to overcome host resistance.
- The discovery of a potential new race with a distinct evolutionary origin highlights the pathogen's adaptive potential and stresses the urgent need for **ongoing disease** surveillance and breeding for novel resistance.

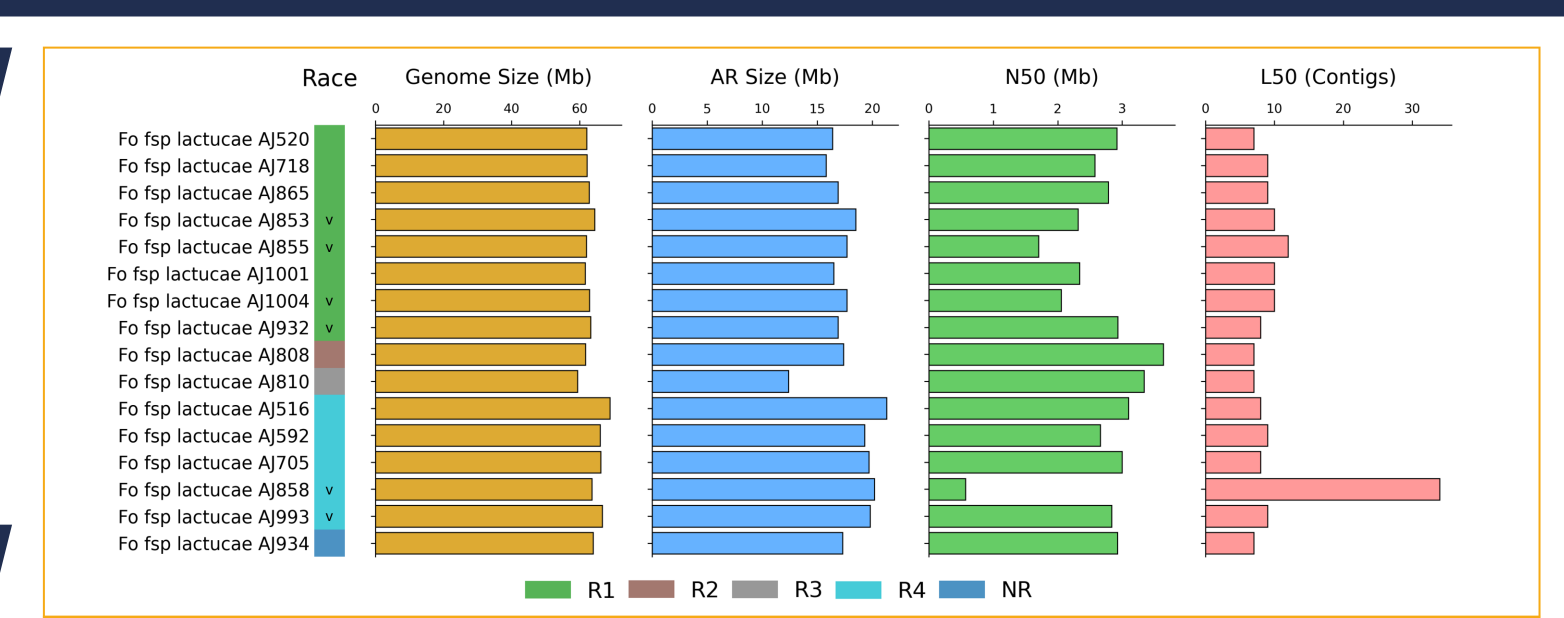


Figure 1. Genome and Accessory Region (AR) sizes, and Assembly Metrics.

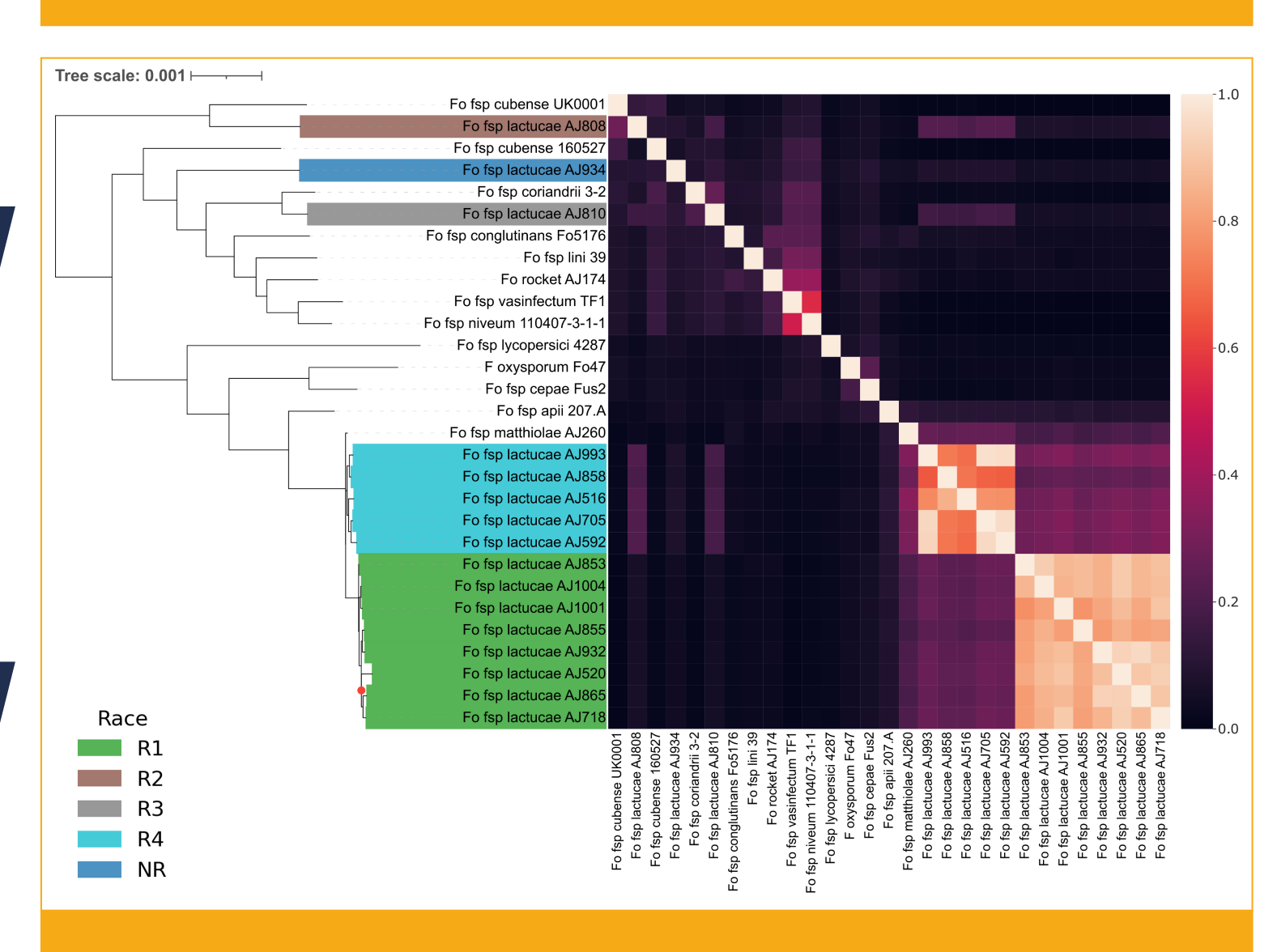


Figure 2. Single Copy Ortholog Phylogeny and AR Similarity Heatmap.

Comparative genomics of the lettuce wilt pathogen reveals a potential new race with a unique evolutionary origin and effector profile.

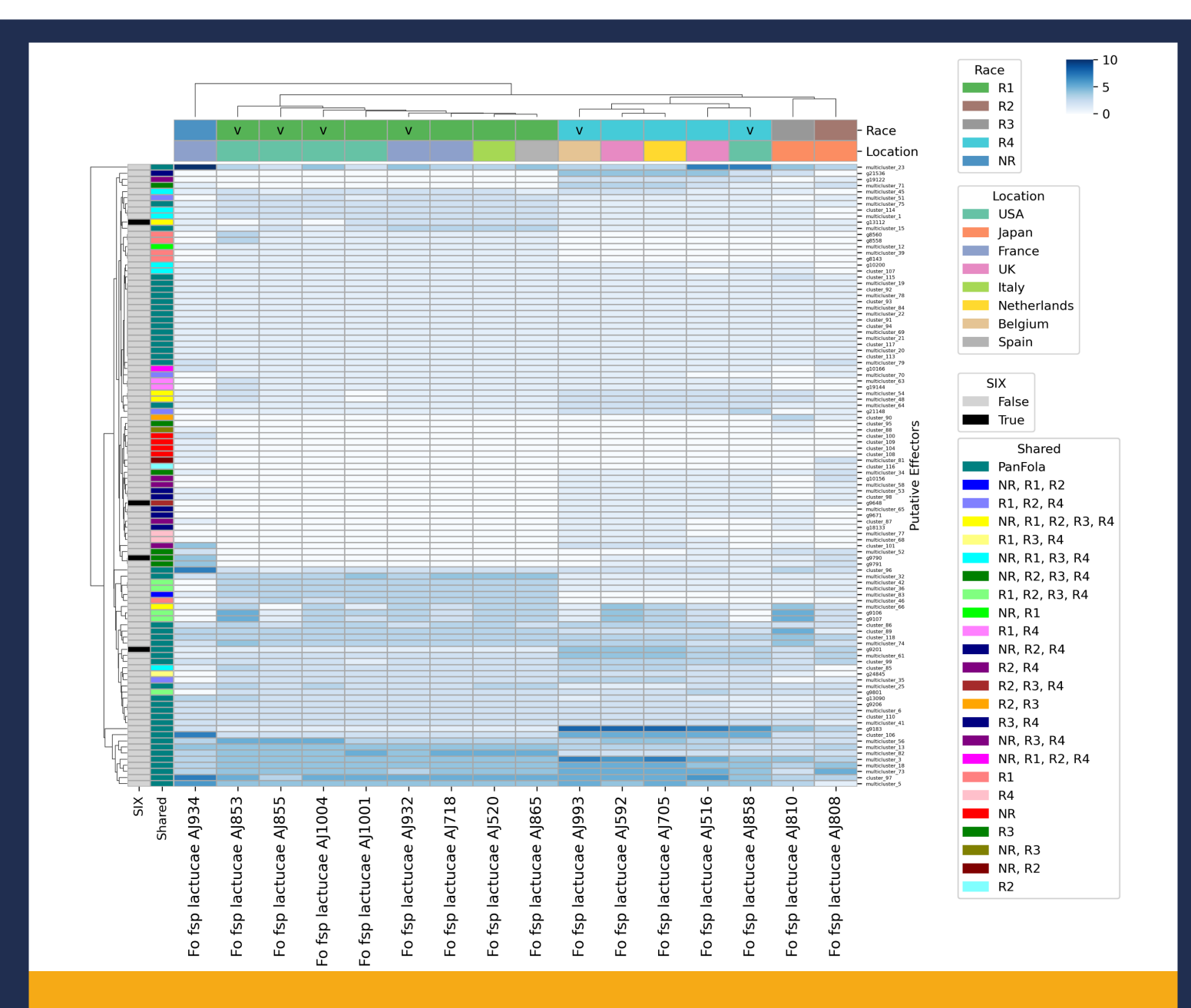


Figure 3. Putative Effector Analysis Heatmap.

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References

- 1. Bates et al., 2024
- 2. van Westerhoven et al., 2024
- 3. Brenes Guallar et al., 2022
- 4. Mestdagh et al. 2025



