ONCLUSIO

NSERVATION GENOMÉ ENDANGERED ALPINE ICHTHYOFAUNA

METHODS

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described pike.

he recent discovery of a pike species, Esox flaviae or Esox cisalpinus² ("Italian pike") native to the Italian region has fuelled debate on the conservation status of this prized game fish and apex predator. Introgressive hybridization with allochthonous Northern pike (E. lucius, here "European pike") is one of the main threats to the genetic identity of the newly

e aim to understand the underlying genomic adaptions of this species to its environment, its population structure and evolutionary relationship with European pike, through a Whole Genome Sequencing (WGS) approach.

Variant calling



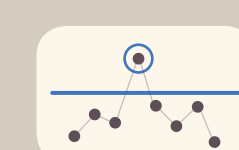
- WGS data at 19x coverage
- 28 samples³: Italian (16), European (6), Hybrid (6)
- GATK Best Practices⁴ yielded 7.3 million SNPs

Population structure



- Principal Component Analysis
- 133k SNPs in Linkage Equilibrium
- Identification of clusters and exclusion of hybrids

Genome-Wide selection scan



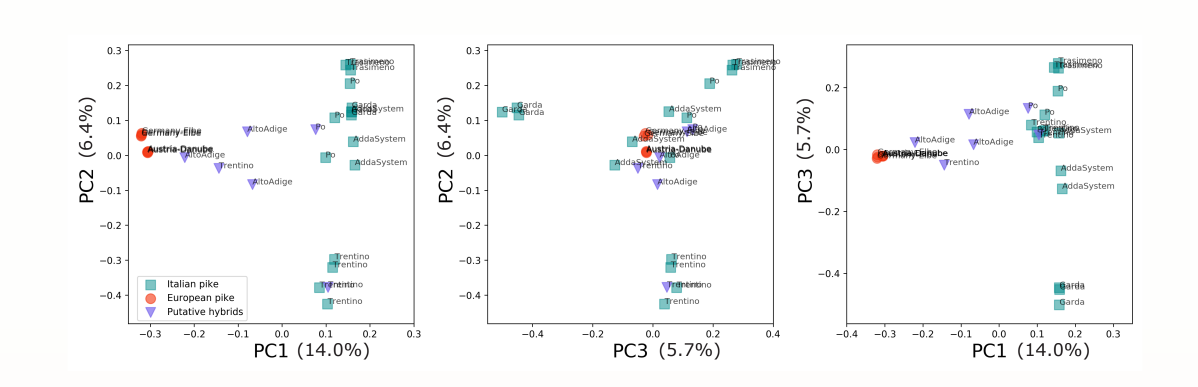
- Genome-wide scans of selection
- Interspecies F_{st} and intraspecies Tajima's D
- 50 Kbp non-overlapping sliding windows

Gene Ontology Enrichment Analysis



- Detection of genes within 50 Kb of outlier F_{st} regions
- GOEA of this gene set using g:Profiler⁵

POPULATION STRUCTURE

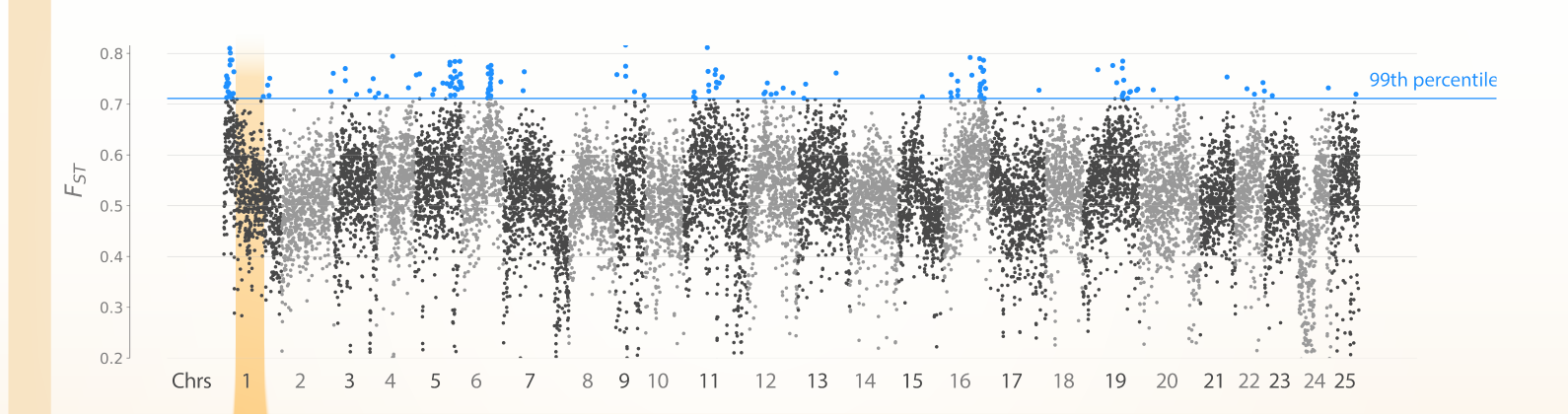


The two species are clearly differentiated in PC1 (14% of variance explained). Also, some differentiation along PC2 (6.4%) and PC3 (5.7%) is present within Italian pike subpopulations, corresponding to the different localities of the samples.

GENOME-WIDE SELECTION SCAN

Candidate chromosomal regions (blue points) under positive selection were detected between species by estimating F_{st} values in non-overlapping 50 Kb windows and selecting outliers above the 99th percentile of the empirical distribution.

Within 50 Kb of these outlier regions, a total of 746 candidate genes were identified.



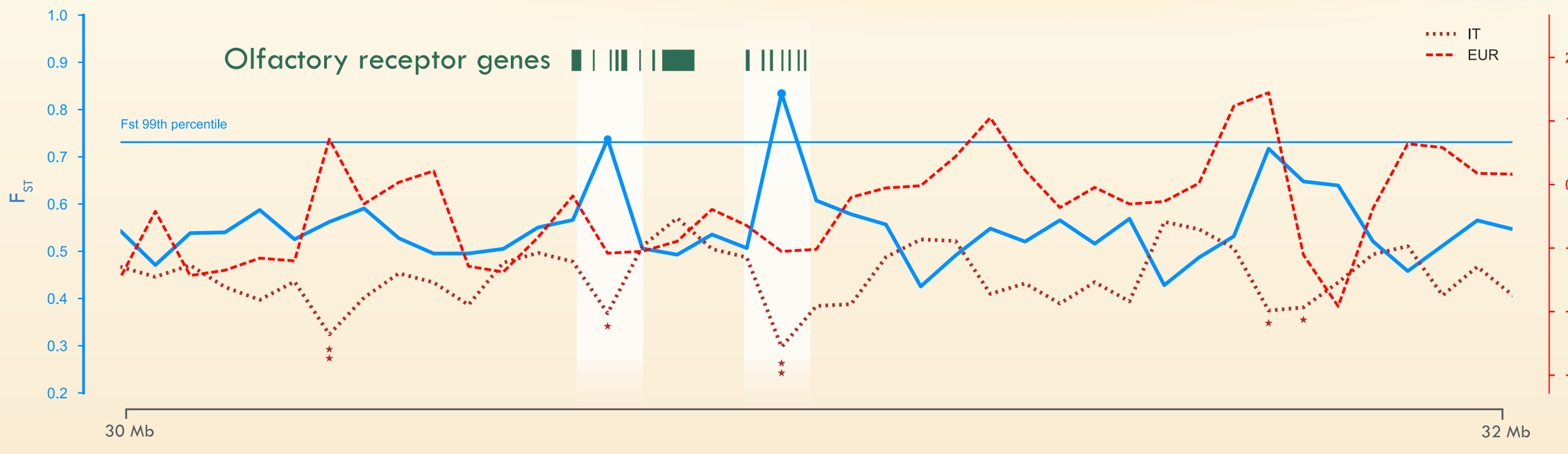
GO ENRICHMENT ANALYSIS

GO ID	GO TERM NAME	ADJUSTED P-VALUE
GO:0050911	Detection of chemical stimulus involved in sensory perception of smell	6.0 x 10 ⁻⁵
GO:0050907	Detection of chemical stimulus involved in sensory perception	7.0 x 10 ⁻⁵
GO:0007608	Sensory perception of smell	8.2 x 10 ⁻⁵
GO:0009593	Detection of chemical stimulus	8.2 x 10 ⁻⁵

Allochthonous European pike

Endemic Italian pike





* and ** respectively indicate Tajima's D negative outliers at the 0.5th and 1st percentiles of the genome-wide empirical distributions. Negative D values suggest positive selection.

Gene Ontology Enrichment Analysis reveals several functional categories significantly enriched for olfactory perception, all attributed to a cluster of 16 G protein-coupled odorant receptor genes on chromosome 1. Within these regions, negative outlier Tajima's D values in Italian but not in European pike suggest positive selection acting in the endemic cisalpine species.

ike, being apex ambush predators, rely greatly on their olfactory perception, perhaps more so than previously believed⁷. Our results suggest olfactory receptors are undergoing positive selection in Italian pike, possibly due to adaptation to local ecosystems. Because this species is threatened by introgressive hybridization with non-native European pike, used in stocking practices for angling purposes, this study informs the need to preserve endemic biodiversity and its genomic adaptations.

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