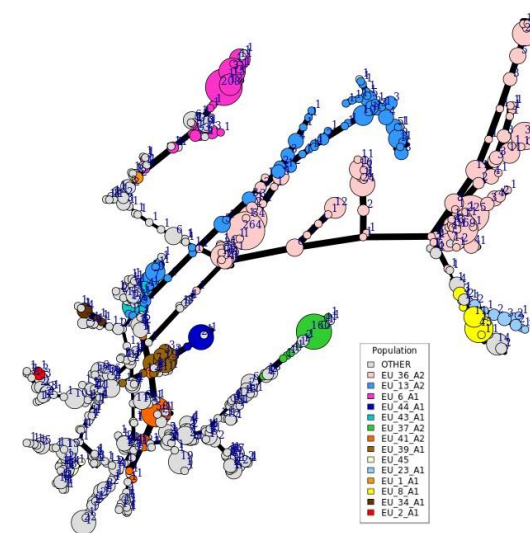




Global early warning for new and emerging *P. infestans* genotypes
Step 1. The *P. infestans* global genetic landscape initiative (PiGGL)
Jens G. Hansen & David Cooke



Time	Speaker/s	Topic
5 mins	Coordination Committee	Introductions
15 mins	David Cooke	Proposal development updates (PiGGL): Jens Hansen or David Cooke present proposal updates. Q&A
10 mins	Zhenxin Zhang	Plans for AsiaBlight: Shipping of FTA cards Zhenxin Zhang provides an update for the cards. Provide info on how to send the cards back. Q&A
25 mins	Md Rashidul Islam	AsiaBlight-Bangladesh updates: Bio-fungicides and Updates on PLB in Bangladesh Q&A
25 mins	Wallace Chen	AsiaBlight-Taiwan updates: Overview of WorldVeg late blight activities Q&A
15 mins	Louise Cooke	EAPR meeting highlights Q&A
5 mins	Coordination Committee	Preparation for the next meeting: <ol style="list-style-type: none"> 1. Select speakers and date (suggested speakers: Seishi Akino and Georgia or Indonesia) 2. Thanks for the reports from members (Saltanat Mambetova) 3. Website - updates and corrections? (including Nepal Regional Representatives)

Late blight early Warning – new genotype/s appears!

Short term

Alerts and transboundary warnings
Farmers adapt IPM strategy accordingly



Effective system for disease surveillance and sampling of isolates (Hunting the new and the rare)

Rapid and reliable characterization of the pathogen

Effective and coordinated communication and dissemination of results (SMS, SoMe, maps & charts, newsletters)

Long term

National list of varieties updated
Breeding programs adapted
Dissemination of results



Assessment of the epidemic potential of new emerging races

Accelerated breeding for resistance

Adaptation of IPM based prevention and control strategies

How can we as a global network contribute?

Late blight early Warning – new genotypes appears!

Short term

Alerts and transboundary warnings
Farmers adapt IPM strategy accordingly



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Assessment of the epidemic potential of new emerging races

Accelerated breeding for resistance

Adaptation of IPM based prevention and control strategies

How will we as a global network contribute?

Indicate the weather based risk via national DSS and Regional risk maps. Data on crop emergence plus crop growth model

Use BlightTracker APP or similar for disease surveillance and sampling. Show data on a Dashboard.

Early sampling - send to national or regional labs. Get results in a week including interpretation. Inform the farming community and other relevant stakeholders

After quality control send data to one central database to build the global picture

Hunt the new in trap nurseries, breeding nurseries and farmers fields, and send samples for genotyping and phenotyping at a regional scale. Interact with breeders and share information. Include the Agrochemical industry in fungicide resistance testing

Test and share best practice re IPM and precision agriculture

Global early warning for new and emerging *P. infestans* genotypes

Super goal: Improved preparedness for new and emerging genotypes or races of *Phytophthora infestans*

Step 1. Activity: “The *P. infestans* global genetic landscape initiative (PiGGL)”

Goal: Generating the *P. infestans* global genetic landscape – to be prepared for new and emerging genotypes of *P. infestans* on a global scale.

- 2022: Preparation, alignment studies, establishing protocols etc.
- 2023: sampling and analysis of 1000-2000 isolates per network (EuroBlight, USABlight, Tizon Latino, AfricaBlight and AsiaBlight).

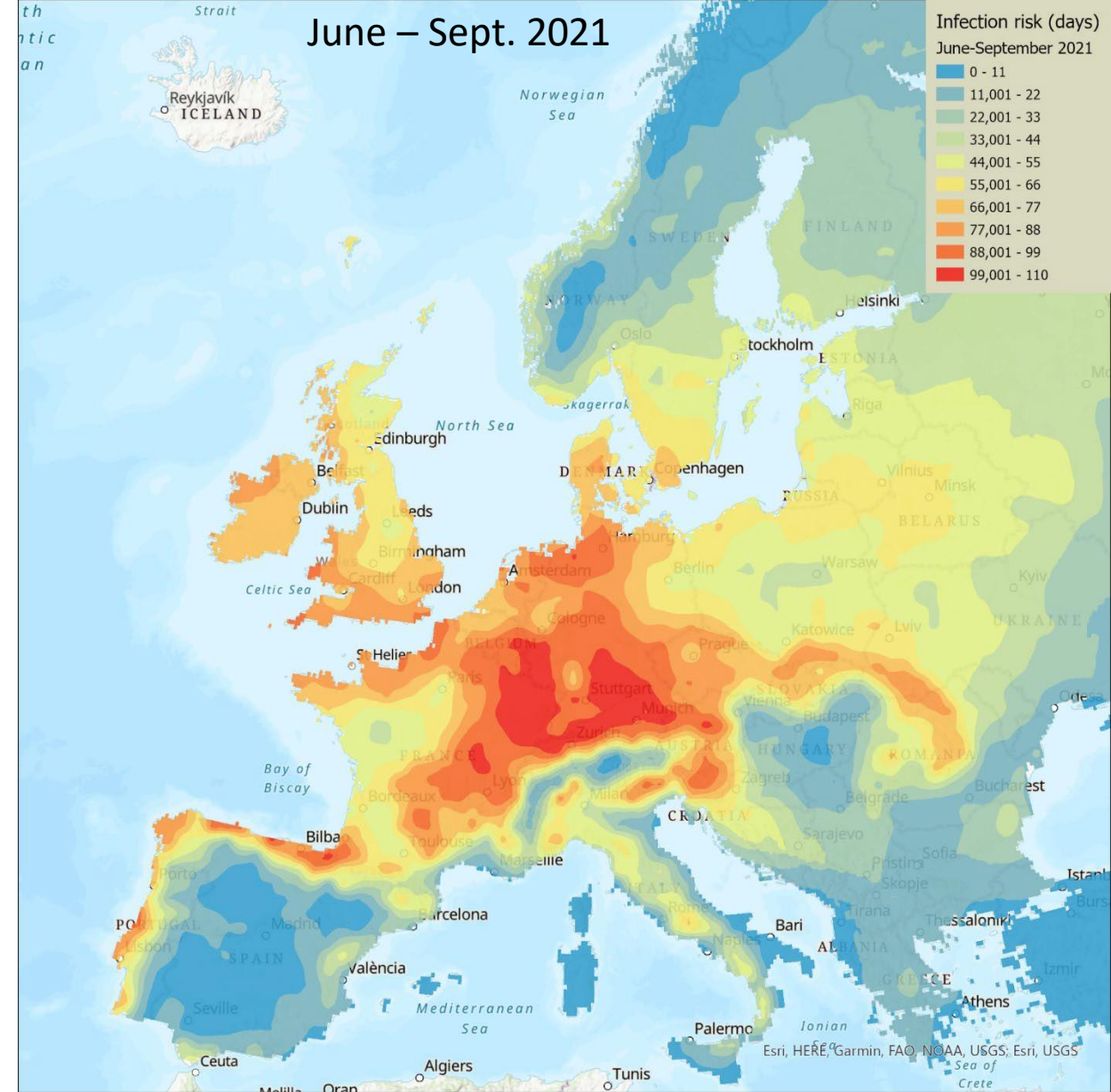
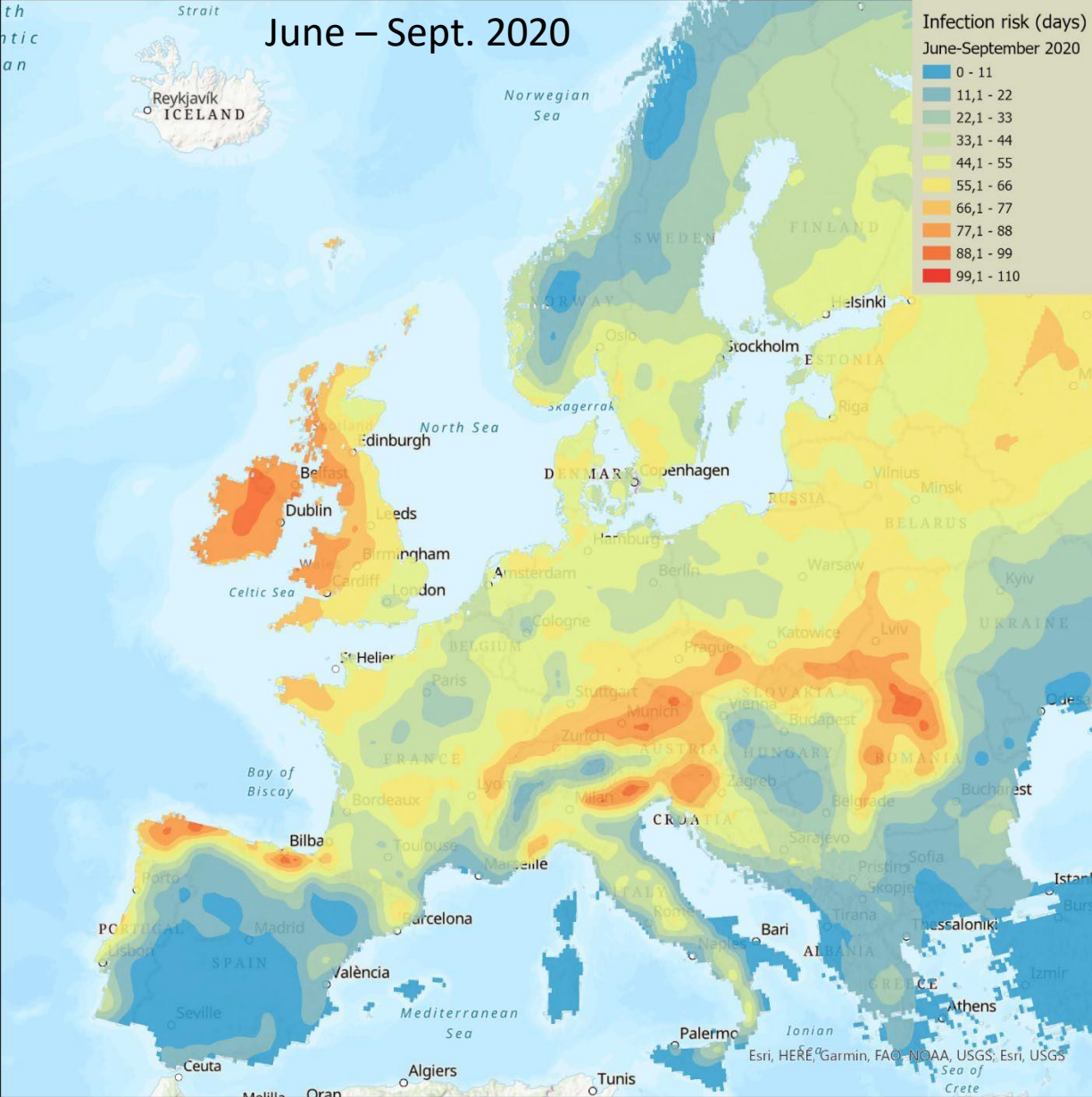
Target activities

Need to have

- Establish and document a global network of dedicated labs covering all 5 PLB continental networks. Labs participate in 2022 in alignment studies and preparation of protocols for sampling and analysis of sampled data. This is ongoing
- In 2023 sample up to 2000 isolates per network, analyse the data with SSR/SNP/Amplicon sequencing or any molecular method agreed upon in the Lab technical group and upload the results into the EuroBlight database
- Publish a scientific paper about the initiative and the results

Nice to have

- Global phenotyping strategy / Platforms - to characterise / phenotype the new genotypes
- Global network of Trap & Field Nurseries - for monitoring resistance in time and space



Identify Late blight disease development hot spots and centres of diversities

GENOTYPE

Genotype map

Genotype frequency map

Genotype frequency chart

Frequency rank

World map

Continent

Europe

Year

- All
- 2021 2020
- 2019 2018
- 2017 2016
- 2015 2014
- 2013 2012
- 2011 2010
- 2009 2008
- 2007 2006
- 2005 2004

Host

- All
- N/A Other
- Potato Tomato

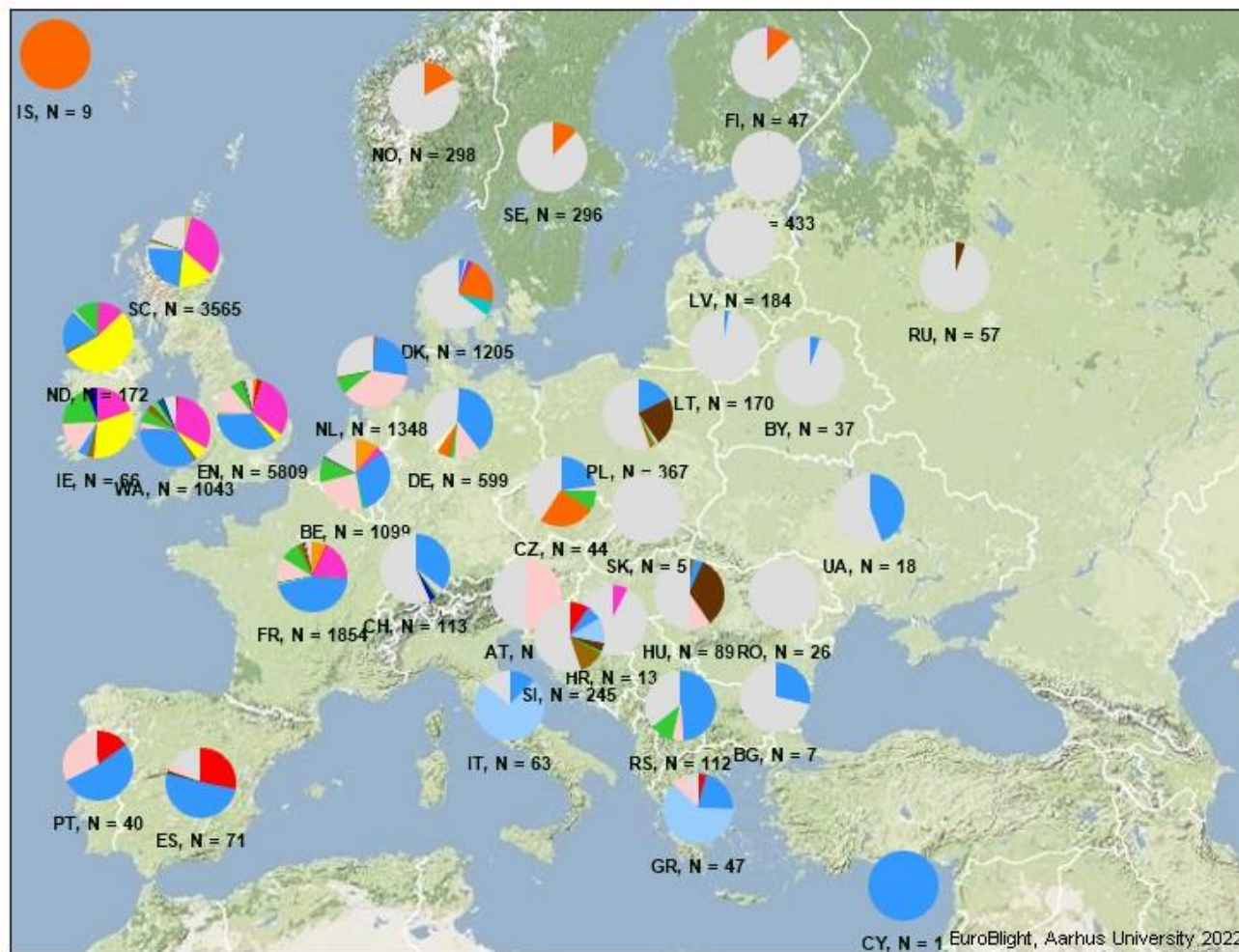
Show

Genotype legend

- | | |
|-----------------------------------|-----------------------------------|
| <input type="checkbox"/> EU_1_A1 | <input type="checkbox"/> EU_2_A1 |
| <input type="checkbox"/> EU_5_A1 | <input type="checkbox"/> EU_6_A1 |
| <input type="checkbox"/> EU_8_A1 | <input type="checkbox"/> EU_12_A1 |
| <input type="checkbox"/> EU_13_A2 | <input type="checkbox"/> EU_23_A1 |
| <input type="checkbox"/> EU_33_A2 | <input type="checkbox"/> EU_34_A1 |
| <input type="checkbox"/> EU_35_A2 | <input type="checkbox"/> EU_36_A2 |
| <input type="checkbox"/> EU_37_A2 | <input type="checkbox"/> SIB_1_A1 |
| <input type="checkbox"/> EU_39_A1 | <input type="checkbox"/> EU_38_A2 |
| <input type="checkbox"/> EU_40_A2 | <input type="checkbox"/> EU_41_A2 |
| <input type="checkbox"/> EU_10_A2 | <input type="checkbox"/> EU_22_A2 |
| <input type="checkbox"/> EU_3_A2 | <input type="checkbox"/> EU_43_A1 |
| <input type="checkbox"/> EU_42_A2 | <input type="checkbox"/> EU_45 |
| <input type="checkbox"/> EU_44_A1 | <input type="checkbox"/> Other |

Genotype frequency distribution

Help



2021: 2495
All years: 42446

Help

2021: 0
All years: 727

Continent
Africa & West/Central Asia

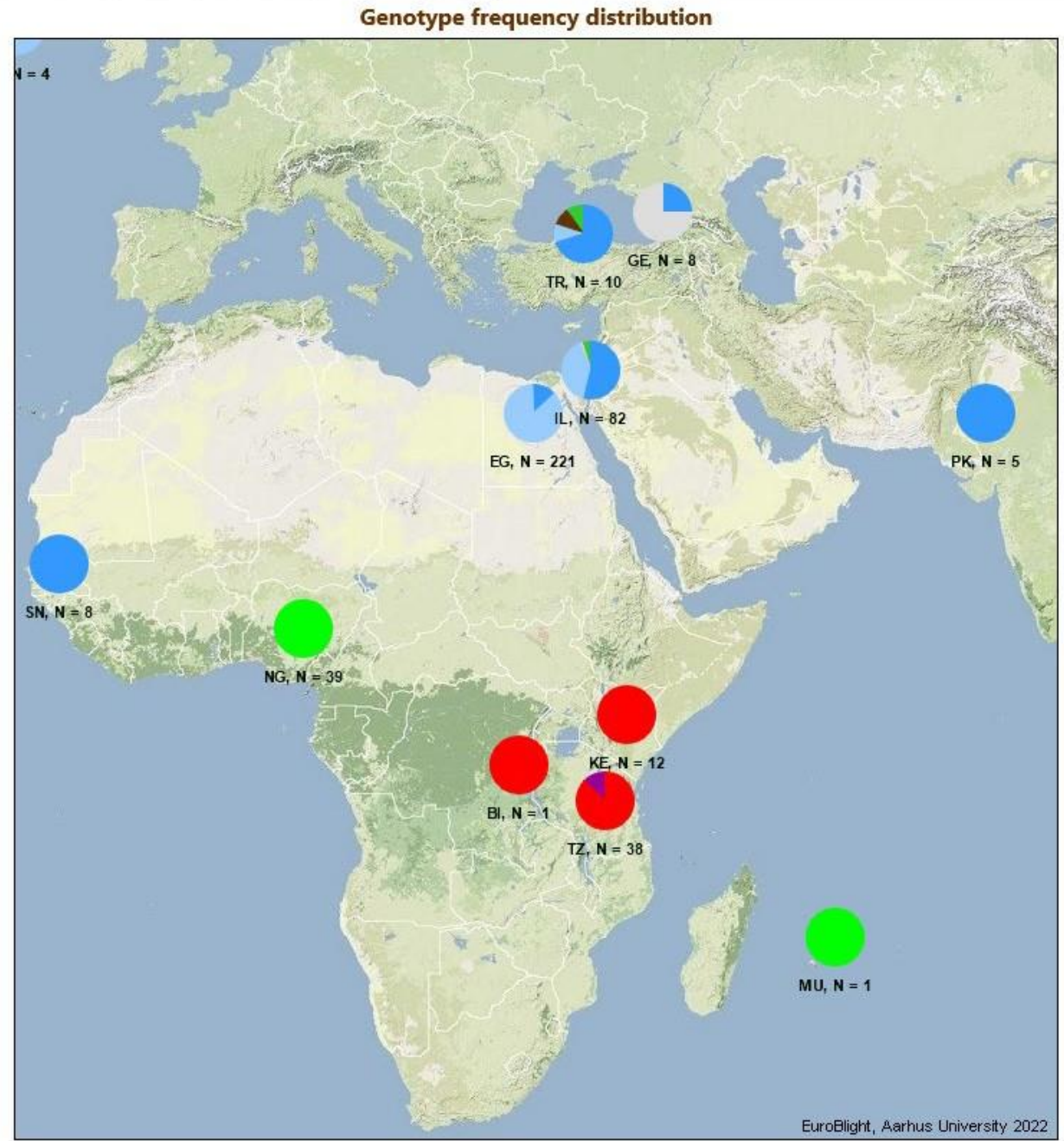
Year
 All
 2020 2019
 2018 2017
 2016 2015
 2014 2013
 2012 2011
 2010

Host
 All
 Potato Tomato

Show

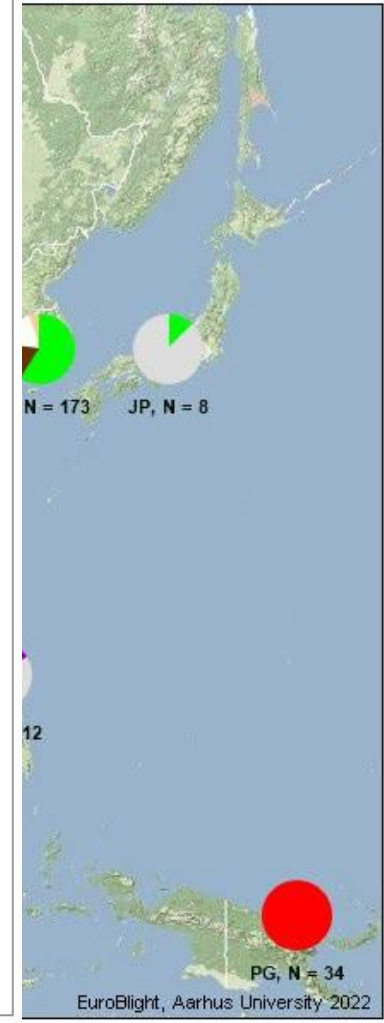
Genotype legend

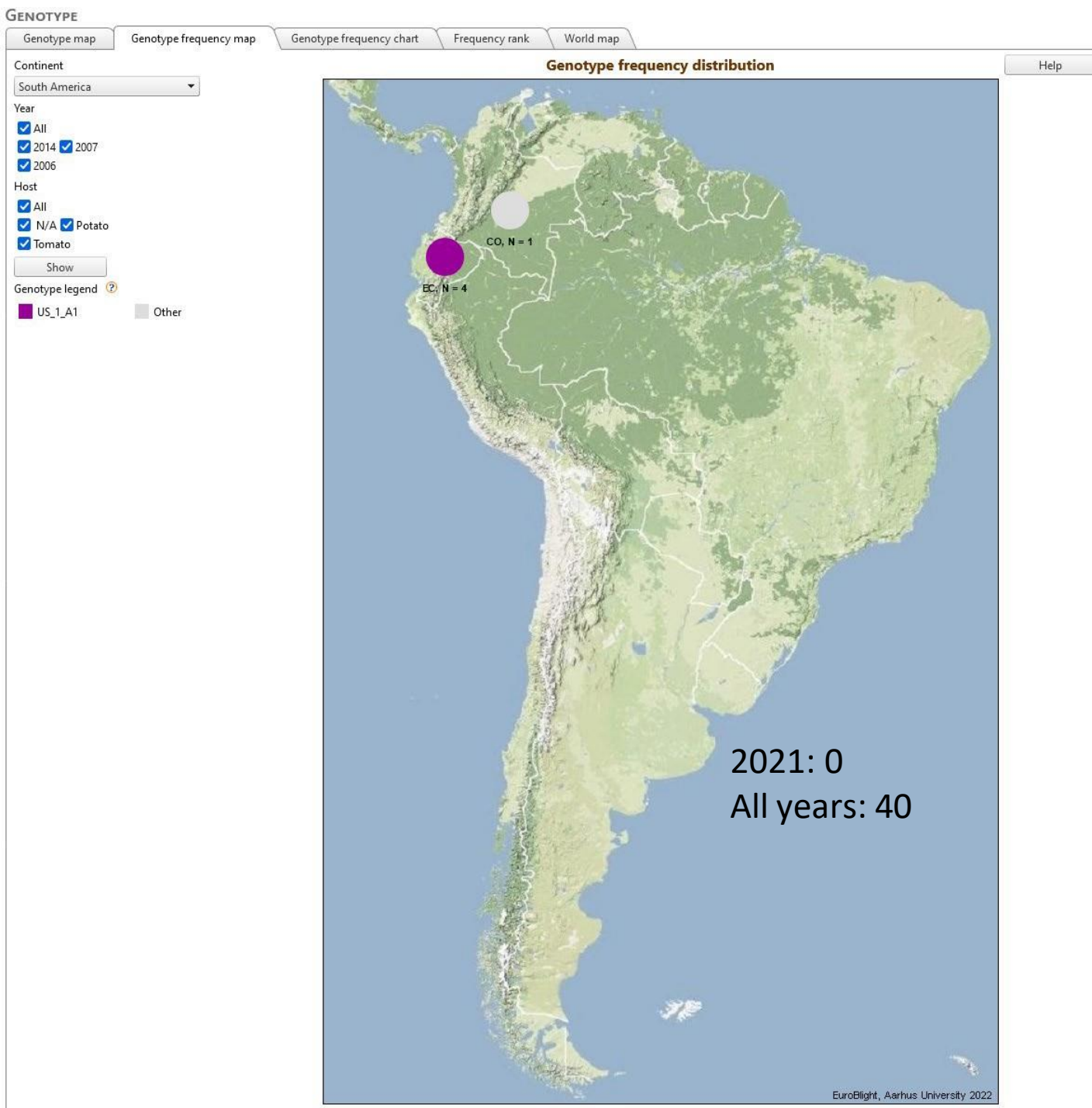
■ EU_2_A1	■ EU_13_A2
■ EU_23_A1	■ EU_33_A2
■ EU_34_A1	■ EU_36_A2
■ EU_37_A2	■ US_1_A1
■ Other	



2021: 0
All years: 885

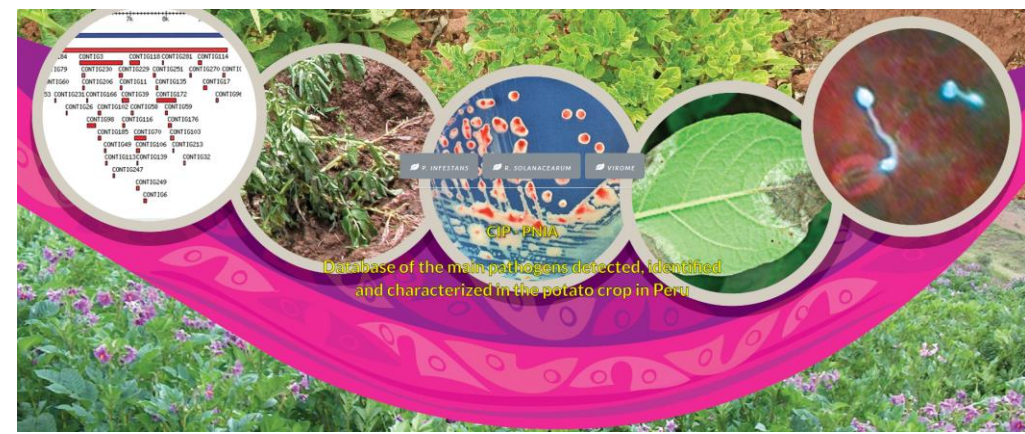
Help





2021: 0
All years: 183

A lot more data to enter from TizonLatino contacts



CIP database

<http://potpathodiv.org/index.html>

Region	Number of Isolates in the EuroBlight database	Period
Europe	42446	1995-2021
Africa & West & Central Asia	727	2008-2020
South/East Asia	885	2005-2019
Oceania	85	2020-12 & 2017
South America	183	2014-2019
North and Central America	40	2016 & 2019

Genotyping centres co-ordinated with EuroBlight standard protocols and scoring



Continent

Europe

Country

All countries selected

Host

- All
- N/A Other Potato

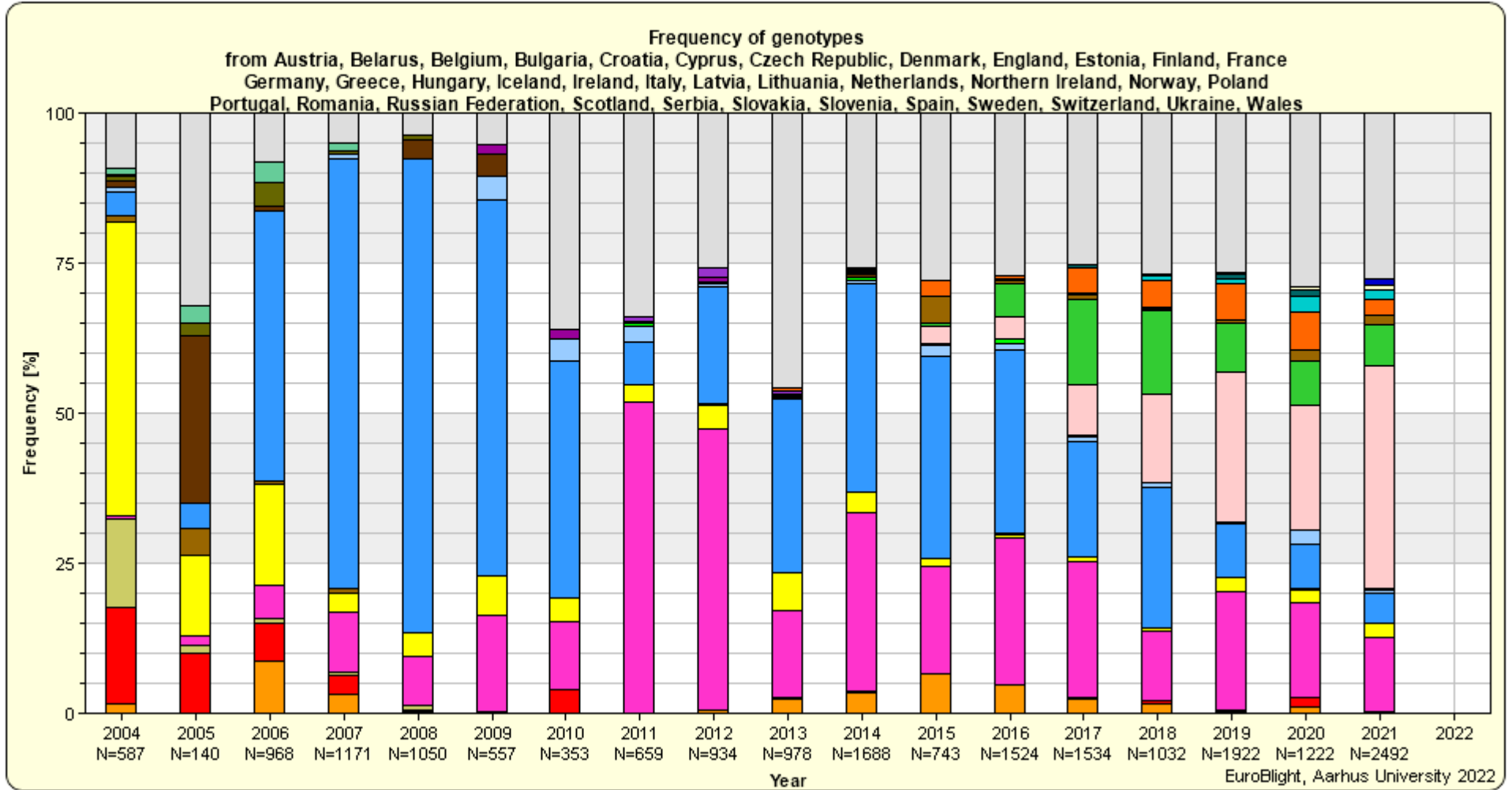
Tomato

Show

Genotype legend ?

- | | |
|--|---|
| ■ EU_1_A1 | ■ EU_2_A1 |
| ■ EU_5_A1 | ■ EU_6_A1 |
| ■ EU_8_A1 | ■ EU_12_A1 |
| ■ EU_13_A2 | ■ EU_23_A1 |
| ■ EU_33_A2 | ■ EU_34_A1 |
| ■ EU_35_A2 | ■ EU_36_A2 |
| ■ EU_37_A2 | ■ SIB_1_A1 |
| ■ EU_39_A1 | ■ EU_38_A2 |
| ■ EU_40_A2 | ■ EU_41_A2 |
| ■ EU_10_A2 | ■ EU_22_A2 |
| ■ EU_3_A2 | ■ EU_43_A1 |
| ■ EU_42_A2 | ■ EU_45 |
| ■ EU_44_A1 | ■ Other |

Genotype frequency distribution





YELLOW RUST GENETIC GROUP OUTPUT

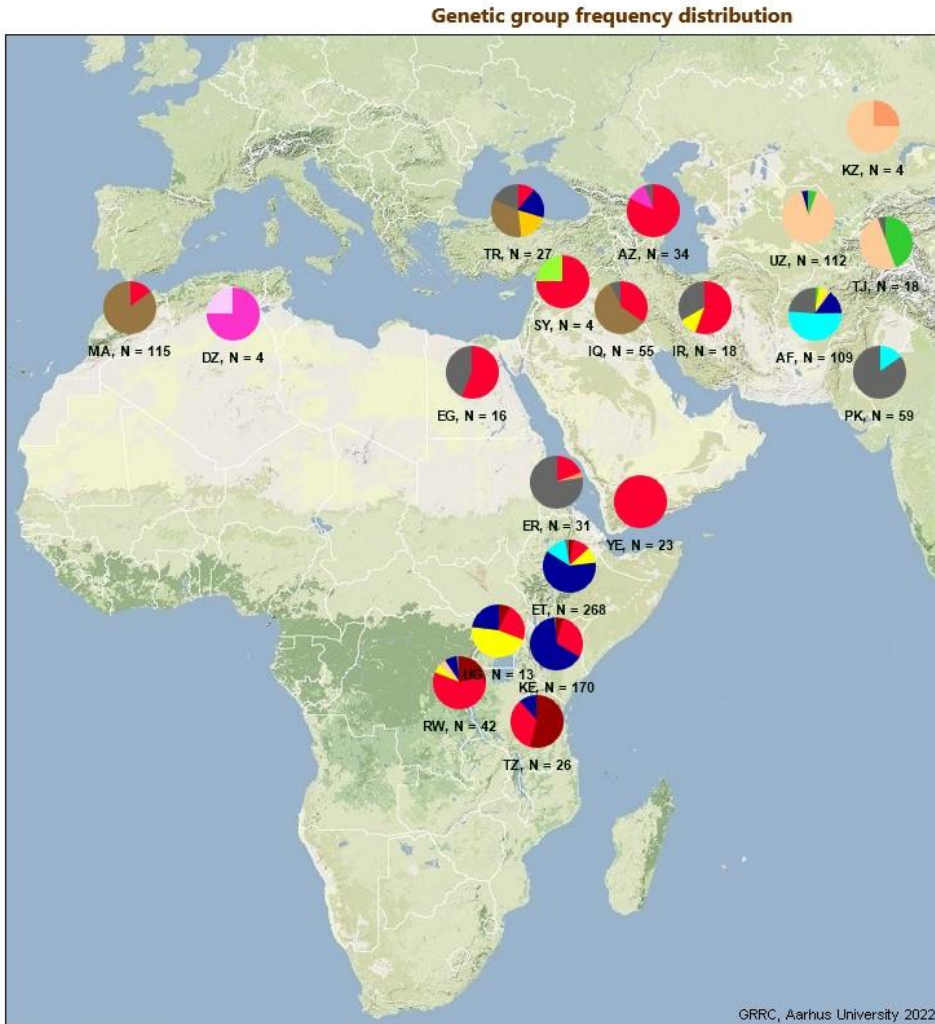
Number of years 19 Width Toolbox GRRC Language

Map Frequency map Frequency chart Frequency rank World map First appearance

Continent
Africa & West/Central Asia
Laboratory
All laboratories selected

Analysis
 SSR genotypes
Year
 All
 2022
 2021
 2020
 2019
 2018
 2017
 2016
 2015
 2014
 2013
 2012
 2011
 2010
 2009
 2007
 2005
 2004

Genetic group
 PstS1
 PstS2
 PstS3
 PstS4
 PstS5
 PstS6
 PstS7
 PstS9
 PstS10
 PstS11
 PstS13
 PstS14
 PstS16
 Other



GRRC, Aarhus University 2022

Data provider: GRRC, Denmark

Bill and Melinda Gates Foundations has funded global wheat rust early warning since 2008

Since 2010 uses the same global database at AU as for *P. infestans*!

The BMGF Foundation has far not supported research and development in Potato

We can change that – if we come up with good arguments!!

Collaborate with the USAID innovation lab Coordinated by David Hughes, Pen state University (CIP included also)

New BMGF proposal on “global infrastructure for transboundary diseases”

P. Infestans will be proposed as a use-case in 2023

Technologies for higher resolution genotyping *P. infestans*

Goal: Extending SSR fingerprinting to sequence-based analyses

WHAT and WHY?

- Harness power of high-throughput and affordable sequencing
- Analysis of more loci in genome than 12 SSRs
- Provide raw data to understand the way genome is evolving in response to selection pressure
 - Mutation rate across different parts of genome
 - Ploidy advantage and destabilisation of genome over time – LOH
- A closer link to phenotype than current neutral SSR markers
 - Effectors – links to R-gene stability
 - Fungicide resistance genes – some targets known and others could be discovered
- Harness network to understand phylogeography and pathogen evolution over time and in space

HOW?

- Baseline SSR typing to identify populations/samples to target
- Whole Genome – 250 Mbases of highly repetitive sequence – very difficult to assemble and process
- Targeted
 - PENSEQ – enrichment of genes of choice followed by Illumina sequence - Aman poster
 - Multiplex effector PCR followed by Illumina sequence Kurt Lamour Uni Tennessee
 - Oxford NanoPore – real-time selective sequencing of key targets (portable?)
- Discussions underway on a phase of pilot studies to compare methods and develop pipelines for data analysis

Roadmap



2022:

- SSR baseline globally and identification of target areas by regional network
- Global coordination group, regional partners and labs identified and displayed on our websites
- Use of molecular technologies identified for genotyping as well as phenotyping
- Ring test study and pipelines for data analysis and display
- Sampling protocol and methodologies applied / genotyping, effector diversity, phenotyping
- Workflow and data flow identified and agreed upon
- Funding of activities by region
- FAIRification, data sharing, IPR, storage, display and publications agreed upon

Discussion at Regional level autumn 2022. Common meeting for all networks in December 2022

2023

Describe the initiative on all network websites and upload protocols

Organise FTA cards and (bar) coding system by network

Conduct regional meetings

Meet in global group in March to update sampling protocol and organisation



Thank you for your attention