Wind-Dispersed Nomadic Diseases: Conclusions for Disease Resistance

E. LIMPERT1* and P. BARTOŠ2

1Phytopathology, Institute of Plant Sciences, Swiss Federal Institute of Technology Zurich (ETH), 8057 Zurich, Switzerland; 2Division of Genetics and Plant Breeding – Research Institute of Crop Production, 161 06 Prague-Ruzyně, Czech Republic
*e-mail: eckhard_limpert@hotmail.com

Abstract: Data on the distribution of some wind-dispersed nomadic plant diseases (powdery mildew, leaf rust) suggest that virulence genes accumulate in the direction of prevailing winds, through the interaction of migration and selection. E.g., for leaf rust of wheat virulence complexity increased from 2–4 on average in France to approximately 7 in Austria and Poland up to 14 and above in Siberia. This situation can be of considerable importance for the use of host resistance. The higher is the virulence in a pathogen population, the more valuable should be the effective resistance genes present in that area for resistance breeding. If virulence complexity increases in the direction of predominant dispersal, the number of unnecessary virulence genes increases as well. The time of efficient use of resistance genes is thought to be prolonged if the area of host culture would shift against the direction of predominant dispersal.

Keywords: nomadic pathogens, cereal diseases, resistance genes, avirulence genes, population biology, wind dispersal, gene flow

An efficient and environmentally benign way of plant protection is through the use of genes conferring disease resistance (R-genes) in the host (PINK & HAND 2002; POLÁK & BARTOŠ 2002). The economic effect, however, varies according to the level of virulence genes (V-genes) enabling the pathogen to overcome the host resistance. For wind-dispersed nomadic diseases a novel pattern of the distribution of V-genes and virulence complexity, VC, i.e. the number of V-genes per pathogen genotype, became evident recently. The evidence suggests that virulence genes accumulate in the direction of prevailing winds, through the interaction of migration and selection. This is the case and, from the underlying logic, appears to have to be so, for pathogens that depend on dispersal to survive, i.e. for obligate nomads, in particular if polycyclic diseases are concerned like the cereal rusts and mildews that are so highly mobile (LIMPERT et al. 2002).

For leaf rust on wheat, VC increased from 2–4 on average in France to approximately 7 in Austria and Poland up to 14 and above in Siberia (after PARK & FELSENSTEIN 1998, re-evaluated; MESTERHÁZY et al. 2000). These and further results from other data sets confirm consistent evidence obtained with the barley mildew pathogen from where such observations and the concept originate (LIMPERT et al. 1999, 2000, and unpublished data).

The concept appears to be significant for any area with prevailing dispersal in certain directions, for continental Europe as well as for Great Britain, where predominant dispersal is from Scotland to England (LIMPERT et al. 2000). The example of Mlo virulence being most expressed in Japan (LYNGKJAER et al. 1995) at the very end of the Eurasian wind path suggests that genes acting quantitatively may also be concerned.

As major conditions of host presence and prevailing westerly winds are similar or the same in Europe and Asia, VC is thought to increase along the Eurasian path of population genetics from western Europe up to eastern Asia (LIMPERT et al. 2002). According to a more novel, fundamental and less established way of thinking it is the number of avirulence genes decreasing in that direction (DE WIT 1992; ROHE et al. 1995; JOOSTEN & DE WIT 1999) but, for convenience, the established terminology is kept here.

CONCLUSIONS AND OUTLOOK

The hypothesis can be of considerable importance for the use of host resistance. The higher the virulence in a pathogen population, the more valuable should be the effective R-genes present in that area. Again, as further
discussed elsewhere (LIMPERT et al. 2000, 2002), this has to be regarded as a general trend and further points affecting the potential value of resistance genes of course continue to need careful consideration.

For example, a wild relative of barley, *Hordeum spontaneum*, is known for its resistance diversity that correlates with population size (JAHOOR & FISCHBECK 1987; DREISEITL & DINOOR 2003). However if, with other conditions being the same, host populations of similar diversity would be compared, originating in Central Europe or the Near East on one hand or in the Far East on the other, from the above hypothesis there should be no doubt that the population from the Far East should be more valuable for disease resistance. Likewise, resistance from the Far East should be more durable according to reasons discussed elsewhere (LIMPERT et al. 1999).

If VC increases in the direction of predominant dispersal, the number of unnecessary virulence genes increases as well. Thus, the hypothesis also casts new light on the discussion of the costs of unnecessary virulence genes, and whether they may or may not be linked with genes for other traits of importance for pathogen fitness.

The present knowledge of the origin of certain R-genes of interest against the wheat powdery mildew pathogen (Table 1) is in line with our reasoning. Almost exclusively, the genes described originate from various regions in Asia, from the Hindukush, India and China. Moreover, in Chinese land races of wheat a number of further genes is expected (HSAM – pers. commun.). The concept should be of interest for other host-pathogen systems involving nomadic pathogens and crops of economic importance such as sugar beet, potato and tobacco, for diseases of grassland, and forest diseases.

Despite the considerable evidence and logic in favour of the hypothesis, it is but too obvious that we only recognised the tip of a novel iceberg so far, where the entire volume would be worth discovering. But one consequence of practical importance for disease resistance relates to the use of host resistance in space. The time of efficient use of R-genes is thought to be prolonged if the area of host culture would shift against the direction of predominant dispersal, rather than in that direction (LIMPERT & BARTOŠ 1997).

For discovering the hypothesis, the cereal pathogens obviously had a pilot function. The hypothesis should be of similar importance for human and animal health, and for further populations affected by wind such as species of insects, spiders, and even some crustaceae living in temporary waters like certain Ostracodae the eggs of which appear to be as easily wind-dispersed as are mildew spores. Therefore, we recently submitted an Expression of Interest for the 6th EU Framework Programme entitled Effects of Wind and Bio-Aerosols on Health and Growth Across Europe, to which further thoughts are always welcome.

**Acknowledgements:** We are grateful to COST at Berne and Brussels for financial support.

### References


### Table 1. Origin of R-genes against wheat powdery mildew (HSAM et al. 2001 for further references, and pers. commun.)

<table>
<thead>
<tr>
<th>Gene</th>
<th>Alleles/Origin</th>
<th>Geographic origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pm3</td>
<td>10 alleles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>a Hope (Australia)</td>
<td>Israel?</td>
</tr>
<tr>
<td></td>
<td>b Mli (Germany)</td>
<td>Hindukush?</td>
</tr>
<tr>
<td></td>
<td>c India</td>
<td></td>
</tr>
<tr>
<td></td>
<td>d spelt?</td>
<td>Asia</td>
</tr>
<tr>
<td></td>
<td>e China</td>
<td></td>
</tr>
<tr>
<td>Pm24</td>
<td></td>
<td>China</td>
</tr>
</tbody>
</table>


