A population-level invasion by transposable elements triggers genome expansion in a fungal pathogen

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- 30 Running title: Transposable element invasion triggers genome expansion

31 ABSTRACT

32 Genome evolution is driven by the activity of transposable elements (TEs). The spread of TEs can 33 have deleterious effects including the destabilization of genome integrity and expansions. However, 34 the precise triggers of genome expansions remain poorly understood because genome size evolution 35 is typically investigated only among deeply divergent lineages. Here, we use a large population 36 genomics dataset of 284 individuals from populations across the globe of Zymoseptoria tritici, a 37 major fungal wheat pathogen. We built a robust map of genome-wide TE insertions and deletions to 38 track a total of 2,456 polymorphic loci within the species. We show that purifying selection 39 substantially depressed TE frequencies in most populations but some rare TEs have recently risen in 40 frequency and likely confer benefits. We found that specific TE families have undergone a 41 substantial genome-wide expansion from the pathogen's center of origin to more recently founded 42 populations. The most dramatic increase in TE insertions occurred between a pair of North 43 American populations collected in the same field at an interval of 25 years. We find that both 44 genome-wide counts of TE insertions and genome size have increased with colonization bottlenecks. 45 Hence, the demographic history likely played a major role in shaping genome evolution within the 46 species. We show that both the activation of specific TEs and relaxed purifying selection underpin 47 this incipient expansion of the genome. Our study establishes a model to recapitulate TE-driven 48 genome evolution over deeper evolutionary timescales.

50 **INTRODUCTION**

51 Transposable elements (TEs) are mobile repetitive DNA sequences with the ability to independently 52 insert into new regions of the genome. TEs are major drivers of genome instability and epigenetic 53 change (Eichler & Sankoff, 2003). Insertion of TEs can disrupt coding sequences, trigger 54 chromosomal rearrangements, or alter expression profiles of adjacent genes (Lim, 1988; Petrov et 55 al., 2003; Slotkin & Martienssen, 2007; Hollister & Gaut, 2009; Oliver et al., 2013). Hence, TE 56 activity can have phenotypic consequences and impact host fitness. While TE insertion dynamics are 57 driven by the selfish interest for proliferation, the impact on the host can range from beneficial to 58 highly deleterious. The most dramatic examples of TE insertions underpinned rapid adaptation of 59 populations or species (Feschotte, 2008; Chuong et al., 2017), particularly following environmental 60 change or colonization events. Beneficial TE insertions are expected to experience strong positive 61 selection and rapid fixation in populations. However, most TE insertions have neutral or deleterious 62 effects upon insertions. Purifying selection is expected to rapidly eliminate deleterious insertions 63 from populations unless constrained by genetic drift (Walser et al., 2006; Baucom et al., 2008; 64 Cridland et al., 2013; Stuart et al., 2016; Lai et al., 2017; Stritt et al., 2017). Additionally, genomic 65 defense mechanisms can disable transposition activity. Across eukaryotes, epigenetic silencing is a 66 shared defense mechanism against TEs (Slotkin & Martienssen, 2007). Fungi evolved an additional 67 and highly specific defense system introducing repeat-induced point (RIP) mutations into any nearly 68 identical set of sequences. The relative importance of demography, selection and genomic defenses 69 determining the fate of TEs in populations remain poorly understood.

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A crucial property predicting the invasion success of TEs in a genome is the transposition rate. TEs tend to expand through family-specific bursts of transposition followed by prolonged phases of transposition inactivity. Bursts of insertions of different retrotransposon families were observed across eukaryotic lineages including *Homo sapiens, Zea mays, Oryza sativa* and *Blumeria graminis* (Shen *et al.*, 1991; SanMiguel *et al.*, 1998; Eichler & Sankoff, 2003; Piegu *et al.*, 2006; Lu *et al.*, 2017; Frantzeskakis *et al.*, 2018). Prolonged bursts without effective counter-selection are thought to

77 underpin genome expansions. In the symbiotic fungus Cenococcum geophilum, the burst of TEs 78 resulted in a dramatically expanded genome compared to closely related species (Peter et al., 2016). 79 Similarly, a burst of a TE family in brown hydras led to an approximately three-fold increase of the 80 genome size compared to related hydras (Wong et al., 2019). Across the tree of life, genome sizes 81 vary by orders of magnitude and enlarged genomes invariably show hallmarks of historic TE 82 invasions (Kidwell, 2002). Population size variation is among the few correlates of genome size 83 across major groups, suggesting that the efficacy of selection plays an important role in controlling 84 TE activity (Lynch, 2007). Reduced selection efficacy against deleterious TE insertions is expected 85 to lead to a ratchet-like increase in genome size. In fungi, TE-rich genomes often show an isochore 86 structure alternating gene-rich and TE-rich compartments (Rouxel et al., 2011). TE-rich 87 compartments often harbor rapidly evolving genes such as effector genes in pathogens or resistance 88 genes in plants (Raffaele & Kamoun, 2012; Jiao & Schneeberger, 2019). Taken together, incipient 89 genome expansions are likely driven by population-level TE insertion dynamics.

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91 The fungal wheat pathogen Zymoseptoria tritici is one of the most important pathogens on crops, 92 causing high yield losses in many years (Torriani et al., 2015). Z. tritici emerged during the 93 domestication of wheat in the Fertile Crescent where the species retained high levels of genetic 94 variation (Zhan et al., 2005; Stukenbrock et al., 2011). The pathogen migrated to all temperate zones 95 where wheat is currently grown and underwent multiple migration bottlenecks, in particular when 96 colonizing Oceania and North America (Zhan et al., 2005; Estep et al., 2015). The genome is 97 completely assembled and shows size variation between individuals sampled across the global 98 distribution range (Feurtey et al., 2020; Badet et al., 2020) (Goodwin et al., 2011). The TE content 99 of the genome shows a striking variation of 17-24% variation among individuals (Badet et al., 100 2020). Z. tritici recently gained major TE-mediated adaptations to colonize host plants and tolerate 101 environmental stress (Omrane et al., 2015, 2017; Krishnan et al., 2018; Meile et al., 2018). Clusters 102 of TEs are often associated with genes encoding important pathogenicity functions (*i.e.* effectors), 103 recent gene gains or losses (Hartmann & Croll, 2017), and major chromosomal rearrangements 104 (Croll et al., 2013; Plissonneau et al., 2016). Transposition activity of TEs also had a genome-wide

105	impact on gene expression profiles during infection (Fouché et al., 2019). The well-characterized
106	demographic history of the pathogen and evidence for recent TE-mediated adaptations make Z.
107	tritici an ideal model to recapitulate the process of TE insertion dynamics, adaptive evolution and
108	changes in genome size at the population level.

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Here, we retrace the population-level context of TE insertion dynamics and genome size changes across the species range by analyzing populations sampled on four continents for a total of 284 genomes. We developed a robust pipeline to detect newly inserted TEs using short read sequencing datasets. Combining analyses of selection and knowledge of the colonization history of the pathogen, we tested whether population bottlenecks were associated with substantial changes in the TE content and the size of genomes.

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118 **Results**

119 A DYNAMIC TE LANDSCAPE SHAPED BY STRONG PURIFYING SELECTION

120 We detected 4,753 TE copies, grouped into 30 families with highly variable copy numbers in the 121 reference genome IPO323 (Figure 2 - figure supplement 1 and Figure 2 - figure supplement 2A). To 122 establish a comprehensive picture of within-species TE dynamics, we analyzed 295 genomes from a 123 worldwide set of six populations spanning the distribution range of the wheat pathogen Z. tritici. To 124 ascertain the presence or absence of TEs across the genome, we developed a robust pipeline (Figure 125 1A). In summary, we called TE insertions by identifying reads mapping both to a TE sequence and a 126 specific location in the reference genome. Then, we assessed the minimum sequencing coverage to 127 reliably recover TE insertions and removed 11 genomes with an average read depth below 15X 128 (Figure 1B). We tested for evidence of TEs using read depth at target site duplications (Figure 1C) 129 and scanned the genome for mapped reads indicating gaps at TE loci (Figure 1D). We found robust 130 evidence for a total of 18,864 TE insertions grouping into 2,465 individual loci. Of these loci, 35.5% 131 (n = 876) have singleton TEs (*i.e.*, this locus is only present in one isolate: Figure 2A, figure

132 supplement 3). An overwhelming proportion of loci (2,345 loci or 95.1%) have a TE frequency 133 below 1%. Singleton TE insertions in particular can be the product of spurious Illumina read 134 mapping errors (Nakamura et al., 2011). To assess the reliability of the detected singletons, we 135 focused on seven isolates for which PacBio long-read data was available (Badet et al., 2020). 136 Aligned PacBio reads confirmed the exact location of 71% (22 out of 31 singleton insertions among 137 seven isolates; see Methods for further details). We found no significant difference in read coverage 138 between confirmed and unconfirmed singleton insertions (Figure 2 - figure supplement 2C,-B and 139 Figure 2 - figure supplement 4).

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141 The abundance of singleton TE insertions strongly supports the idea that TEs actively copy into new 142 locations but also indicates that strong purifying selection maintains nearly all TEs at low frequency 143 (Figure 2A). The density of TE loci on accessory chromosomes, which are not shared among all 144 isolates of the species, is almost twice the density found on core chromosomes (102 versus 58 TEs 145 per Mb; Figure 2B and Figure 2 - figure supplement 5A). This suggests relaxed selection against TE 146 insertion on the functionally dispensable and gene-poor accessory chromosomes. We found no 147 difference in TE allele frequency distribution between recombination hotspots and the rest of the 148 genome (Figure 2 - figure supplement 5B). Similarly, the TE density and the number of insertions 149 did not vary between recombination hotspots and the genomic background (Figure 2 - figure 150 supplement 5C).

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TEs grouped into 23 families and 11 superfamilies, with 88.2% of all copies belonging to class I/retrotransposons (n = 2175; Figure 2C and Figure 2 - figure supplements 6A-B). RLG/*Gypsy* (n =1,483) and RLC/*Copia* (n = 623) elements constitute the largest long terminal repeats (LTR) superfamilies. Class II/DNA transposons are dominated by DHH/*Helitron* (n = 249). As expected, TE families shared among fewer isolates tend to show also lower global copy numbers (*i.e.*, all isolates combined), while TE families that are present in all isolates generally have high global copy numbers (Figure 2D).

160 We detected 153 loci with TEs inserted into genes with most of the insertions being singletons 161 (44.7%; n = 68) or of very low frequency (Figure 2E). Overall, TE insertions into exonic sequences 162 were less frequent than expected compared to insertions into up- and downstream regions, which is 163 consistent with effective purifying selection (Figure 2F). Insertions into introns were also strongly 164 under-represented, likely due to the small size of most fungal introns (~ 50-100 bp) and the high 165 probability of disrupting splicing or adjacent coding sequences. We also found that insertions 800-166 1000 bp away from coding sequences of a focal gene were under-represented. Given the high gene 167 density, with an average spacing between genes of 1.744 kb, TE insertions within 800-1,000 bp of a 168 coding gene tend to be near adjacent genes already. Taken together, TEs in the species show a high 169 degree of transposition activity and are subject to strong purifying selection.

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171 DETECTION OF CANDIDATE TE LOCI UNDERLYING RECENT ADAPTATION

172 The TE transposition activity can generate adaptive genetic variation. To identify the most likely 173 candidate loci, we analyzed insertion frequency variation among populations as an indicator for 174 recent selection. Across all populations, the insertion frequencies differed only weakly with a strong 175 skew towards extremely low F_{ST} values (mean = 0.0163; Figure 3A-B and Figure 3 - figure 176 supplement 1). To further analyze evidence for TE-mediated adaptive evolution, we screened a 177 genome-wide SNP dataset for evidence of selective sweeps using selection scans. We found 16.5 % 178 of all TE loci located in regions of selective sweep. Given our population sampling of two 179 population pairs, we tested for adaptive TE insertions in selective sweep regions either in the North 180 American or European population pairs. Hence, we selected loci having low TE insertion 181 frequencies (< 5%) in all populations except either the recent North American or European 182 population (> 20%) (Figure 3B). Based on these criteria, we obtained 7 candidate loci possibly 183 underlying local adaptation (6 in North America, one in Europe; Figure 4A and Figure 4 - figure 184 supplement 1). All loci carry inserted retrotransposons with 4 RLG Luna, one RLG Mercurius and 185 one RLG_Deimos.

187 One TE insertion is 3,815 bp downstream of a gene encoding an RTA1-like protein, which can 188 function as transporters with a transmembrane domain and have been associated with resistance 189 against several antifungal compounds (Soustre et al., 1996). The insertion is also 5785 bp upstream 190 of a gene encoding a protein kinase domain (Figure 4B). The TE insertion was not detected in the 191 Middle East or the two European populations, and was at low frequencies in the Australian (3.7%) 192 and North American 1990 (1.7%) populations, but increased to 53% of all isolates in the North 193 American 2015 population (fixation index $F_{ST} = 0.42$; Figure 4 - figure supplement 1). Isolates that 194 carry the insertion show a significantly higher resistance to azole antifungal compounds (Figure 4C). 195 The TE is in the subtelomeric region of chromosome 12, with a moderate GC content, a low TE and 196 a high gene density (Figure 4D). The TE belongs to the family RLG Luna, which shows a 197 substantial burst across different chromosomes within the species (Figures 4E-F). We found no 198 association between the phylogenetic relationships among isolates based on the two closest genes 199 and the presence or absence of the TE insertion (Figure 4G). A second candidate adaptive TE 200 insertion belongs to the RLG_Mercurius family and is located between two genes of unknown 201 function (Figure 4 - figure supplement 2). A third potentially adaptive TE insertion of a 202 RLC Deimos is 229 bp upstream of a gene encoding a SNARE domain protein and 286 bp upstream 203 of a gene encoding a flavin amine oxidoreductase. Furthermore, the TE is inserted in a selective 204 sweep region (Figure 4 - figure supplement 2). SNARE domains play a role in vesicular transport 205 and membrane fusion (Bonifacino & Glick, 2004). An additional four candidates for adaptive TE 206 insertions belong to RLG Luna and were located distantly to genes (Figure 4 - figure supplement 2). 207 We experimentally tested whether the TE insertions in proximity to genes were associated with 208 higher levels of fungicide resistance. For this, we measured growth rates of the fungal isolates in the 209 presence or absence of an azole fungicide widely deployed against the pathogen. We found that the 210 insertion of TEs at two loci was positively associated with higher levels of fungicide resistance, 211 suggesting that the adaptation was mediated by the TE (Figure 4C and Figure 4 - figure supplement 212 2).

214 POPULATION-LEVEL EXPANSIONS IN TE CONTENT

215 If TE insertion dynamics are largely neutral across populations, TE frequencies across loci should 216 reflect neutral population structure. To test this, we performed a principal component analysis based 217 on a set of six populations on four continents that represent the global genetic diversity of the 218 pathogen (Figure 5A) and 900,193 genome-wide SNPs (Figure 5B). The population structure 219 reflected the demographic history of the pathogen with clear continental differentiation and only 220 minor within-site differentiation. To account for the lower number of TE loci, we performed an 221 additional principal component analysis using a random SNP set of similar size to the number of TE 222 loci. The reduced SNP set retained the geographic signal of the broader set of SNPs (Figure 5C). In 223 stark contrast, TE frequencies across loci showed only weak clustering by geographic origin with the 224 Australian population being the most distinct (Figure 5D). We found a surprisingly strong 225 differentiation of the two North American populations sampled at a 25-year interval in the same field 226 in Oregon.

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228 Unusual patterns in population differentiation at TE loci suggests that TE activity may substantially 229 vary across populations (Figure 6, Figure 4 - figure supplement 1). To analyze this, we first 230 identified the total TE content across all loci per isolate. We found generally lower TE numbers in 231 the Middle Eastern population from Israel (Figure 6A-C, and Figure 6 - figure supplement 1), which 232 is close to the pathogen's center of origin (Stukenbrock et al., 2007). Populations that underwent at 233 least one migration bottleneck showed a substantial burst of TEs across all major superfamilies. 234 These populations included the two populations from Europe collected in 1999 and 2016 and the 235 North American population from 1990, as well as the Australian population. We found a second 236 stark increase in TE content in the North American population sampled in 2015 at the same site as 237 the population from 1990. Strikingly, the isolate with the lowest number of analyzed TEs collected 238 in 2015 was comparable to the isolate with the highest number of TEs at the same site in 1990. We 239 tested whether sequencing coverage could explain variation in the detected TEs across isolates, but 240 we found no meaningful association (Figure 2 - figure supplement 6C). We analyzed whether the 241 population-specific expansions were correlated with shifts in the frequency spectrum of TEs in the

242 populations (Figure 6D). We found that the first step of expansions observed in Europe compared to 243 the Middle East (Israel) was associated with an upwards shift in allele frequencies. This is consistent 244 with transposition activity creating new copies in the genomes and stronger purifying selection in the 245 Middle East. Similarly, the North American populations showed also signatures consistent with 246 relaxation of selection against TEs (i.e., fewer low frequency TEs). We found a significant 247 difference (Two-sample Kolmogorov-Smirnov test, two-sided) in the curve shapes between the 248 population from the Middle East and North America 2015 (Figure 6 - figure supplement 2). We 249 analyzed variation in TE copy numbers across families and found that the expansions were mostly 250 driven by RLG elements including the families Luna, Sol and Venus, the RLC family Deimos and 251 the LINE family Lucy (Figure 6E and Figure 6 - figure supplement 3A). We also found a North 252 American specific burst in DHH elements of the family Ada (increase from 4.6 to 6.1 copies on 253 average per isolate), an increase specific to Swiss populations in LINE elements, and an increase in 254 RLC elements in the Australian and the two North American populations. Analyses of complete Z. 255 tritici reference-quality genomes that include isolates from the Israel, Australia, Switzerland (1999) 256 and North American (1990) population revealed high TE contents in Australia and North America 257 (Oregon 1990) (Badet et al., 2020). The reference-quality genomes confirmed also that the increase 258 in TEs was driven by LINE, RLG and RLC families in Australia and DHH, RLG and RLC families 259 in North America (Badet et al., 2020).

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261 TE-MEDIATED GENOME SIZE EXPANSIONS

262 The combined effects of actively copying TE families and relaxed purifying selection leads to an 263 accumulation of new TE insertions in populations. Consequently, mean genome sizes in populations 264 should increase over generations. We estimated the cumulative length of TE insertions based on the 265 length of the corresponding TE consensus sequences and found a strong increase in the total TE 266 length in populations outside the Middle East center of origin, and a second increase between the 267 two North American populations (Figure 1 - figure supplement 1). To test for incipient genome 268 expansions within the species, we first assembled genomes of all 284 isolates included in the study. 269 Given the limitations of short-read assemblies, we implemented corrective measures to compensate

270 for potential variation in assembly qualities. We corrected for variation in the GC content of 271 different sequencing datasets by downsampling reads to generate balanced sequencing read sets prior 272 to assembly (see Methods). We also excluded all reads mapping to accessory chromosomes because 273 different isolates are known to differ in the number of these chromosomes. Genome assemblies were 274 checked for completeness by retrieving the phylogenetically conserved BUSCO genes (Figure 7A). 275 Genome assemblies across different populations carry generally >99% complete BUSCO gene sets, 276 matching the completeness of reference-quality genomes of the same species (Badet et al., 2020). 277 The completeness of the assemblies showed no correlation with either TE or GC content of the 278 genomes. GC content was inversely correlated with genome size consistent with the expansion of 279 repetitive regions having generally low GC content (Figure 7B). We found that the core genome size 280 varied substantially among populations with the Middle East, Australia as well as the two older 281 European and North American populations having the smallest core genome sizes (Figure 7C). We 282 found a notable increase in core genome size in both the more recent European and North American 283 populations. The increase in core genome size is positively correlated with the count and cumulative 284 length of all inserted TEs (Figure 7D, 7E and 7G) and negatively correlated with the genome-wide 285 GC content (Figure 7F and 7G). Hence, core genome size shows substantial variation within the 286 species matching the recent expansion in TEs across continents. We found the most variable genome 287 sizes in the more recent North American population (Figure 7 - figure supplement 1B). Finally, we 288 contrasted variation in genome size with the detected TE insertion dynamics. For this, we assessed 289 the variable genome segment as the difference between the smallest and largest analyzed core 290 genome. To reflect TE dynamics, we calculated the cumulative length of all detected TE insertions 291 in any given genome. We found that the cumulative length of inserted TEs represents between 4.8 292 and 184 % of the variable genome segment defined for the species or 0.2-2.6% of the estimated 293 genome size per isolate (Figure 7 - figure supplement 1C-D).

295 **DISCUSSION**

TEs play a crucial role in generating adaptive genetic variation within species but are also drivers of deleterious genome expansions. We analyzed the interplay of TEs with selective and neutral processes including population differentiation and incipient genome expansions. TEs have substantial transposition activity in the genome but are strongly counter-selected and are maintained at low frequency. TE dynamics showed distinct trajectories across populations with more recently established populations having higher TE content and a concurrent expansion of the genome.

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303 RECENT SELECTION ACTING ON TE INSERTIONS

304 TE frequencies in the species show a strong skew towards singleton insertions across populations. 305 However, our short read based analyses are possibly skewed towards over-counting singletons as 306 indicated by independent long-read mapping evaluations. Nevertheless, the skew towards low 307 frequency TE insertions indicates both that TEs are undergoing transposition and that purifying 308 selection maintains frequencies at a low level. Similar effects of selection on active TEs were 309 observed across plants and animals, including Drosophila melanogaster and Brachypodium 310 distachyon (Cridland et al., 2013; Stritt et al., 2017; Luo et al., 2020). TE insertions were under-311 represented in or near coding regions, showing a stronger purifying selection against TEs inserting 312 into genes. Coding sequences in the Z. tritici genome are densely packed with an average distance of 313 only ~1 kb (Goodwin et al., 2011). Consistent with this high gene density, TE insertions were most 314 frequent at a distance of 200-400 bp away from coding sequences. A rapid decay in linkage 315 disequilibrium in the Z. tritici populations (Croll et al., 2015; Hartmann et al., 2018) likely 316 contributed to the efficiency of removing deleterious insertions. Some TE superfamilies have 317 preferred insertion sites in coding regions and transcription start sites (Miyao et al., 2003; Fu et al., 318 2013; Gilly et al., 2014; Quadrana et al., 2016). Hence, some heterogeneity in the observed insertion 319 site distribution across the genome is likely due to insertion preferences of individual TEs. We also 320 found evidence for positive selection acting on TEs with the strongest candidate locus being a TE 321 insertion on chromosome 12. This locus showed a frequency increase only in the more recent North

American population, which experienced the first systematic fungicide applications and subsequent emergence of fungicide resistance in the decade prior to the last sampling (Estep *et al.*, 2015). The nearest gene encodes a RTA1-like protein, a transmembrane exporter which is associated with resistance towards different stressors, including antifungal compounds, and shows strong copy number variation in several fungi (Soustre *et al.*, 1996; Rogers & Barker, 2003; Sirisattha *et al.*, 2004; Ali *et al.*, 2014; Yew *et al.*, 2016; Liang *et al.*, 2018). Hence, the TE insertion may have positively modulated RTA1 expression to resist antifungals.

329 Transposition activity in a genome and counter-acting purifying selection are expected to establish 330 an equilibrium over evolutionary time (Charlesworth & Charlesworth, 1983). However, temporal 331 bursts of TE families and changes in population size due to bottlenecks or founder events are likely 332 to shift the equilibrium. Despite purifying selection, we were able to detect signatures of positive 333 selection by scanning for short-term population frequency shifts. Population genomic datasets can be 334 used to identify the most likely candidate loci underlying recent adaptation. The shallow genome-335 wide differentiation of Z. tritici populations provides a powerful background to test for outlier loci 336 (Hartmann et al., 2018). We found the same TE families to have experienced genome-wide copy 337 number expansions, suggesting that the availability of adaptive TE insertions may be a by-product of 338 TE bursts in individual populations.

339

340 POPULATION-LEVEL TE INVASIONS AND RELAXED SELECTION

341 Across the surveyed populations from four continents, we identified substantial variation in TE 342 counts per genome. The increase in TEs matches the global colonization history of the pathogen 343 with an increase in TE copies in more recently established populations (Zhan et al., 2003; 344 Stukenbrock et al., 2007). Compared to the Israeli population located nearest the center of origin in 345 the Middle East, the European populations showed a three-fold increase in TE counts. The 346 Australian and North American populations established from European descendants retained high 347 TE counts. We identified a second increase at the North American site where TE counts nearly 348 doubled again over a 25-year period. Compared to the broader increase in TEs from the Middle East, 349 the second expansion at the North American site was driven by a small subset of TE families alone.

350 Analyses of completely assembled reference-quality genomes from the same populations confirmed 351 that genome expansions were primarily driven by the same TE families belonging to the RLG, RLC 352 and DHH superfamilies (Badet et al., 2020). Consistent with the contributions from individual TEs, 353 we found that the first expansion in Europe led to an increase in low-frequency variants, suggesting 354 higher transposition activity of many TEs in conjunction with strong purifying selection. The second 355 expansion at the North American site shifted TE frequencies upwards, suggesting relaxed selection 356 against TEs. The population-level context of TEs in Z. tritici shows how heterogeneity in TE control 357 interacts with demography to determine extant levels of TE content and, ultimately, genome size.

358

359 TE INVASION DYNAMICS UNDERPINS GENOME SIZE EXPANSIONS

360 The number of detected TEs was closely correlated with core genome size, hence genome size 361 expansions were at least partly caused by the very recent proliferation of TEs. Genome assemblies of 362 large eukaryotic genomes based on short read sequencing are often fragmented and contain chimeric 363 sequences (Nagarajan & Pop, 2013). Focusing on the less repetitive core chromosomes in the 364 genome of Z. tritici reduces such artefacts substantially. Because genome assemblies are the least 365 complete in the most repetitive regions, any underrepresented sequences may rather underestimate 366 than overestimate within-species variation in genome size. Hence, we consider the assembly sizes to 367 be a robust correlate of total genome size. The core genome size differences observed across the 368 species range match genome size variation typically observed among closely related species. Among 369 primates, genome size varies by ~70% with ~10% between humans and chimpanzees (Rogers & 370 Gibbs, 2014; Miga et al., 2020). In fungi, genome size varies by several orders of magnitude within 371 phyla but is often highly similar among closely related species (Raffaele & Kamoun, 2012). 372 Interestingly, drastic changes in genome size have been observed in the Blumeria and 373 Pseudocercospora genera where genome size changed by 35-130% between the closest known 374 species (González-Sayer et al.; Frantzeskakis et al., 2018). Beyond analyses of TE content variation 375 correlating with genome size evolution, proximate mechanisms driving genome expansions are 376 poorly understood. By establishing large population genetic datasets, such as those possible for crop 377 pathogens, analyses of genome size evolution become tractable at the population level.

378 TEs might not only contribute to genome expansion directly by adding length through additional 379 copies, but also by increasing the rate of chromosomal rearrangements and ectopic recombination 380 (Bourque et al., 2018; Blommaert, 2020). However, TEs are not the only repetitive elements that can 381 lead to a genome size expansion. In Arabidopsis thaliana genomes, the 45S rDNA has been shown 382 to have the strongest impact on genome size variation, followed by 5S rDNA variation, and 383 contributions by centromeric repeats and TEs (Long et al., 2013). In conjunction, recent work 384 demonstrates how repetitive sequences are drivers of genome size evolution over short evolutionary 385 timescales.

386 The activity of TEs is controlled by complex selection regimes within species. Actively transposing 387 elements may accelerate genome evolution and underpin expansions. Hence, genomic defenses 388 should evolve to efficiently target recently active TEs. Here, we show that TE activity and 389 counteracting genomic defenses have established a tenuous equilibrium across the species range. We 390 show that population subdivisions are at the origin of highly differentiated TE content within a 391 species matching genome size changes emerging over the span of only decades and centuries. In 392 conclusion, population-level analyses of genome size can recapitulate genome expansions typically 393 observed across much deeper time scales providing fundamentally new insights into genome 394 evolution.

395

396 **Methods**

397 FUNGAL ISOLATE COLLECTION AND SEQUENCING

We analyzed 295 *Z. tritici* isolates covering six populations originating from four geographic locations and four continents (Figure 5 - figure supplement 1), including: Middle East 1992 (n = 30isolates, Nahal Oz, Israel), Australia 2001 (n = 27, Wagga Wagga), Europe 1999 (n = 33, Berg am Irchel, Switzerland), Europe 2016 (n = 52, Eschikon, ca. 15km from Berg am Irchel, Switzerland), North America 1990 and 2015 (n = 56 and n = 97, Willamette Valley, Oregon, United States) (McDonald *et al.*, 1996; Linde *et al.*, 2002; Zhan *et al.*, 2002, 2003, 2005). Illumina short read data from the Middle Eastern, Australian, European 1999 and North American 1990 populations were

obtained from the NCBI Sequence Read Archive (SRA) under the BioProject PRJNA327615
(Hartmann *et al.*, 2017). For the Switzerland 2016 and Oregon 2015 populations, asexual spores
were harvested from infected wheat leaves from naturally infected fields and grown in YSB liquid
media including 50 mgL⁻¹ kanamycin and stored in silica gel at -80°C. High-quality genomic DNA
was extracted from liquid cultures using the DNeasy Plant Mini Kit from Qiagen (Venlo,
Netherlands). The isolates were sequenced on an Illumina HiSeq in paired-end mode and raw reads
were deposited at the NCBI SRA under the BioProject PRJNA596434.

412

413 TE INSERTION DETECTION

414 The quality of Illumina short reads was determined with FastQC version 0.11.5 415 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) (Figure 1A). To remove spuriously 416 sequenced Illumina adaptors and low quality reads, we trimmed the sequences with Trimmomatic 417 version 0.36, using the following filter parameters: illuminaclip:TruSeq3-PE-2.fa:2:30:10 leading:10 418 trailing:10 slidingwindow:5:10 minlen:50 (Bolger et al., 2014). We created repeat consensus 419 sequences for TE families (sequences are available on https://github.com/crolllab/datasets; Figure 1 -420 figure supplement 5) in the complete reference genome IPO323 (Goodwin et al., 2011) with 421 RepeatModeler version open-4.0.7 (http://www.repeatmasker.org/RepeatModeler/) based on the 422 RepBase Sequence Database and de novo (Bao et al., 2015). TE classification into superfamilies and 423 families was based on an approach combining detection of conserved protein sequences and tools to 424 detect non-autonomous TEs (Badet et al., 2020). To detect TE insertions, we used the R-based tool 425 ngs te mapper version 79ef861f1d52cdd08eb2d51f145223fad0b2363c integrated into the 426 McClintock pipeline version 20cb912497394fabddcdaa175402adacf5130bd1, using bwa version 427 0.7.4-r385 to map Illumina short reads, samtools version 0.1.19 to convert alignment file formats 428 and R version 3.2.3 (Li & Durbin, 2009; Li et al., 2009; Linheiro & Bergman, 2012; R Core Team, 429 2017; Nelson et al., 2017).

431 DOWN-SAMPLING ANALYSIS

432 We performed a down-sampling analysis to estimate the sensitivity of the TE detection with 433 ngs te mapper based on variation in read depth. We selected one isolate per population matching 434 the average coverage of the population. We extracted the per-base pair read depth with the 435 genomecov function of bedtools version 2.27.1 and calculated the genome-wide mean read depth 436 (Quinlan & Hall, 2010). The number of reads in the original fastq file was reduced in steps of 10% 437 to simulate the impact of reduced coverage. We analyzed each of the obtained reduced read subsets 438 with ngs_te_mapper using the same parameters as described above. The correlation between the 439 number of detected insertions and the read depth was visualized using the function nls with model 440 SSlogis in R and visualized with ggplot2 (Wickham, 2016). The number of detected TEs increased 441 with the number of reads until reaching a plateau indicating saturation (Figure 1B). Saturation was 442 reached at a coverage of approximately 15X, hence we retained only isolates with an average read 443 depth above 15X for further analyses. We thus excluded one isolate from the Oregon 2015 444 population and ten isolates from the Switzerland 2016 population.

445

446 VALIDATION PROCEDURE FOR PREDICTED TE INSERTIONS

447 ngs_te_mapper detects the presence but not the absence of a TE at any given locus. We devised 448 additional validation steps to ascertain both the presence as well as the absence of a TE across all 449 loci in all individuals. TEs absent in the reference genome were validated by re-analyzing mapped 450 Illumina reads. Reads spanning both parts of a TE sequence and an adjacent chromosomal sequence 451 should only map to the reference genome sequence and cover the target site duplication of the TE 452 (Figure 1C). We used bowtie2 version 2.3.0 with the parameter --very-sensitive-local to map 453 Illumina short reads of each isolate on the reference genome IPO323 (Langmead & Salzberg, 2012). 454 Mapped Illumina short reads were then sorted and indexed with samtools and the resulting bam file 455 was converted to a bed file with the function bamtobed in bedtools. We extracted all mapped reads 456 with an end point located within 100 bp of the target site duplication (Figure 1C). We tested whether 457 the number of reads with a mapped end around the target site duplication significantly deviated if the 458 mapping ended exactly at the boundary. A mapped read ending exactly at the target site duplication

boundary is indicative of a split read mapping to a TE sequence absent in the reference genome. To test for the deviation in the number of read mappings around the target site duplication, we used a Poisson distribution and the *ppois* function in R version 3.5.1 (Figure 1C). We identified a TE as present in an isolate if tests on either side of the target site duplication had a *p*-value < 0.001 (Figure 5 - figure supplement 1; Figure 1 - figure supplement 1B and Figure 1 - figure supplement 2).

464

465 For TEs present in the reference genome, we analyzed evidence for spliced junction reads spanning 466 the region containing the TE. Spliced reads are indicative of a discontinuous sequence and, hence, 467 absence of the TE in a particular isolate (Figure 1D). We used STAR version 2.5.3a to detect spliced 468 junction reads with the following set of parameters: --runThreadN 1 --outFilterMultimapNmax 100 -469 470 outFilterMismatchNmax 5 --alignIntronMin 150 --alignIntronMax 15000 (Dobin et al., 2012). We 471 then sorted and indexed the resulting bam file with samtools and converted split junction reads with 472 the function bam2hints in bamtools version 2.5.1 (Barnett et al., 2011). We selected loci without 473 overlapping spliced junction reads using the function intersect in bedtools with the parameter -loj -v. 474 We considered a TE as truly absent in an isolate if ngs te mapper did not detect a TE and evidence 475 for spliced junction reads were found, indicating that the isolate had no inserted TE in this region. If 476 the absence of a TE could not be confirmed by spliced junction reads, we labelled the genotype as 477 missing. Finally, we excluded TE loci with more than 20% missing data from further investigations 478 (Figure 1D and Figure 1 - figure supplement 1C).

479

480 CLUSTERING OF TE INSERTIONS INTO LOCI

481 We identified insertions across isolates as being the same locus if all detected TEs belonged to the 482 same TE family and insertion sites differed by ≤ 100 bp (Figure 1 - figure supplement 3). We used 483 the R package GenomicRanges version 1.28.6 with the functions makeGRangesFromDataFrame and 484 findOverlaps and the R package devtools version 1.13.4 (Lawrence et al., 2013; Wickham & Chang, 485 2016). the package dplyr version 0.7.4 summarize We used R to datasets 486 (https://dplyr.tidyverse.org/). Population-specific frequencies of insertions were calculated with the 487 function allele.count in the R package hierfstat version 0.4.22 (Goudet, 2005). We conducted a 488 principal component analysis for TE insertion frequencies filtering for a minor allele frequency \geq 489 5%. We also performed a principal component analysis for genome-wide single nucleotide 490 polymorphism (SNP) data obtained from Hartmann et al (2017) and Singh et al (2020). As described 491 previously, SNPs were hard-filtered with VariantFiltration and SelectVariants tools integrated in the 492 Genome Analysis Toolkit (GATK) (McKenna et al., 2010). SNPs were removed if any of the 493 following filter conditions applied: QUAL<250; QD<20.0; MQ<30.0; -2 > BaseQRankSum > 2; -2 494 > MQRankSum > 2; -2 > ReadPosRankSum > 2; FS>0.1. SNPs were excluded with vcftools version 495 0.1.17 and plink version 1.9 requiring a genotyping rate >90% and a minor allele frequency >5%496 (https://www.cog-genomics.org/plink2, Chang et al., 2015). Finally, we converted tri-allelic SNPs to 497 bi-allelic SNPs by recoding the least frequent allele as a missing genotype. Principal component 498 analysis was performed using the gdsfmt and SNPRelate packages in R (Zheng et al., 2012, 2017). 499 For a second principal component analysis with a reduced set of random markers, we randomly 500 selected SNPs with vcftools and the following set of parameters: --maf 0.05 --thin 200000 to obtain 501 an approximately equivalent number of SNPs as TE loci.

502

503 EVALUATION OF SINGLETON INSERTIONS

504 To evaluate the reliability of singleton TE insertion loci, we analyzed singleton loci in isolates for 505 which we had both Illumina datasets and complete reference-quality genomes (Badet et al., 2020). 506 From a set of 19 long-read PacBio reference genomes spanning the global distribution of Z. tritici, 507 one isolate each from Australia, Israel, North America (1990) and four isolates from Europe (1999) 508 were also included in the TE insertion screening. To assess the reliability of singleton TE insertions, 509 we first investigated structural variation analyses among the reference genomes (Badet *et al.*, 2021, 510 Supplementary Data 1 and 2). The structural variation was called both based on split read mapping 511 of PacBio reads and pairwise whole-genome alignments. Using bedtools intersect, we recovered for 512 the 31 singleton TE loci in the 7 analyzed genomes a total of 17 loci showing either an indel, 513 translocation, copy number polymorphism, duplication, inverted duplication, inversion, or inverted

514 translocation at the same location. We visually inspected the PacBio read alignment bam files 515 against the IPO323 reference genome using IGV version 2.4.16 (Robinson et al., 2011), and found a 516 typical coverage increase at the target site duplication, with most read mappings interrupted at the 517 target site duplication as expected for an inserted TE. For the 14 remaining TE loci, we extracted the 518 region of the predicted insertion and padded the sequence on both ends with an additional 500 bp 519 using samtools faidx. We used blast to identify a homologous region in the assembled reference-520 quality genomes. Matching regions were inspected based on blastn for the presence of a TE 521 sequence matching the TE family originally detected at the locus. With this second approach, we 522 confirmed an additional five singletons to be true insertions. Both methods combined produced 523 supportive evidence for 22 out of 31 singleton insertions (71%). We calculated the read coverage 524 after mapping to the reference genome IPO323 with bedtools genomecov for each PacBio long-read 525 dataset and calculated mean coverage for 500 bp regions around singleton TE insertions.

526

527 POPULATION DIFFERENTIATION IN TE FREQUENCIES

528 We calculated Nei's fixation index (F_{ST}) between pairs of populations using the R packages *hierfstat* 529 and adegenet version 2.1.0 (Jombart, 2008; Jombart & Ahmed, 2011). To understand the 530 chromosomal context of TE insertion loci across isolates, we analyzed draft genome assemblies. We 531 generated *de novo* genome assemblies for all isolates using SPAdes version 3.5.0 with the parameter 532 --careful and a kmer range of "21, 29, 37, 45, 53, 61, 79, 87" (Bankevich et al., 2012). We used 533 blastn to locate genes adjacent to TE insertion loci on genomic scaffolds of each isolate. We then 534 extracted scaffold sequences surrounding 10 kb up- and downstream of the localized gene with the 535 function faidx in samtools and reverse complemented the sequence if needed. Then, we performed 536 multiple sequence alignments for each locus across all isolates with MAFFT version 7.407 with 537 parameter --maxiterate 1000 (Katoh & Standley, 2013). We performed visual inspections to ensure 538 correct alignments across isolates using Jalview version 2.10.5 (Waterhouse et al., 2009). To 539 generate phylogenetic trees of individual gene or TE loci, we extracted specific sections of the 540 alignment using the function extractalign in EMBOSS version 6.6.0 (Rice et al., 2000) and 541 converted the multiple sequence alignment into PHYLIP format with imodeltest version 2.1.10 using

542 the -getPhylip parameter. We then estimated maximum likelihood phylogenetic trees with the 543 software PhyML version 3.0, the K80 substitution model and 100 bootstraps on the ATGC South of 544 France bioinformatics platform (Guindon & Gascuel, 2003; Guindon et al., 2010; Darriba et al., 545 2012). Bifurcations with a supporting value lower than 10% were collapsed in TreeGraph version 546 2.15.0-887 beta and trees were visualized as circular phylograms in Dendroscope version 2.7.4 547 (Huson et al., 2007; Stöver & Müller, 2010). For loci showing complex rearrangements, we 548 generated synteny plots using 19 completely sequenced genomes from the same species using the R 549 package genoplotR version 0.8.9 (Guy et al., 2010; Badet et al., 2020). We calculated the 550 population-specific allele frequency for TE loci and estimated the exponential decay curve with a 551 self-starting Nls asymptomatic regression model nls(p_loci ~ SSasymp(p_round, Asym, R0, lrc) in 552 R.

553 We analyzed signatures of selective sweeps based on genome-wide SNPs using the extended 554 haplotype homozygosity (EHH) tests implemented in the R package REHH (Sabeti et al., 2007; 555 Gautier & Vitalis, 2012). We analyzed within-population signatures based on the iHS statistic and 556 chose a maximum gap distance of 20 kb. We also analyzed cross-population signatures based on the 557 XP-EHH statistic for the following two population pairs: North America 1990 versus North America 558 2015, Europe 1999 versus Europe 2016. We defined significant selective sweeps as being among the 559 99.9th percentile outliers of the iHS and XP-EHH statistics. Significant SNPs at less than 5 kb were 560 clustered into a single selective sweep region adding +/- 2.5 kb. Finally, we analyzed whether TE 561 loci in the population pairs were within 10 kb of a region identified as a selective sweep by XP-EHH 562 using the function intersect from bedtools.

563

564 GENOMIC LOCATION OF TE INSERTIONS

To characterize the genomic environment of TE insertion loci, we split the reference genome into non-overlapping windows of 10 kb using the function splitter from EMBOSS. TEs were located in the reference genome using RepeatMasker providing consensus sequences from RepeatModeler (http://www.repeatmasker.org/). To analyze coding sequence, we retrieved the gene annotation for

569 the reference genome (Grandaubert et al., 2015). We estimated the percentage covered by genes or 570 TEs per window using the function intersect in bedtools. Additionally, we calculated the GC content 571 using the tool (https://github.com/spundhir/RNAget gc content 572 Seq/blob/master/get_gc_content.pl). We extracted the number of TEs present in 1 kb windows 573 around each annotated core gene in the reference genome IPO323, using the function window in 574 bedtools. We calculated the relative distances between each gene and the closest TE with the 575 function bedtools closest. For the TEs inserted into genes, we used the function intersect in bedtools 576 to distinguish intron and exon insertions with the parameters -wo and -v, respectively. TEs that 577 overlap more than one exon were only counted once. For each 100 bp segment in the 1 kb windows 578 as well as for introns and exons, we calculated the mean number of observed TE insertions per base 579 pair. We calculated the mean number of TEs per window and calculated the log2 of the observed 580 number of TE insertions divided by the expected value. We extracted information about 581 recombination hotspots from Croll et al. (2015). This dataset is based on two experimental crosses 582 initiated from isolates included in our analyses (1A5x1E4, 3D1x3D7). The recombination rates were 583 assessed based on the reference genome IPO323 and analyzed with the R/qtl package in R. We used 584 bedtools intersect to compare both TE density in IPO323 and TE insertion polymorphism with 585 predicted recombination hotspots.

586

587 CORE GENOME SIZE ESTIMATION

588 Accessory chromosomes show presence/absence variation within the species and length 589 polymorphism (Goodwin et al., 2011; Croll et al., 2013) and thus impact genome size. We 590 controlled for this effect by first mapping sequencing reads to the reference genome IPO323 using 591 bowtie2 with --very-sensitive-local settings and retained only reads mapping to any of the 13 core 592 chromosomes using seqtk subseq v1.3-r106 (https://github.com/lh3/seqtk/). Furthermore, we found 593 that different sequencing runs showed minor variation in the distribution of the per read GC content. 594 In particular, reads of a GC content lower than 30 % were underrepresented in the Australian (mean 595 reads < 30% of the total readset: 0.05\%), North American 1990 (0.07\%) and Middle East (0.1\%) 596 populations, and higher in the Europe 1999 (1.3 %), North American 2015 (3.0 %) and Europe 2016

(4.02 %) populations (Figure 1 - figure supplement 4). Library preparation protocols and Illumina
sequencer generations are known factors influencing the recovery of reads of varying GC content
(Benjamini & Speed, 2012).

600

601 To control a potential bias stemming from this, we subsampled reads based on GC content to create 602 homogeneous datasets. For this, we first retrieved the mean GC content for each read pair using 603 geecee in EMBOSS and binned reads according to GC content. For the bins with a GC content 604 <30%, we calculated the mean proportion of reads from the genome over all samples. We then used 605 seqtk subseq to subsample reads of <30% to adjust the mean GC content among readsets. We 606 generated *de novo* genome assemblies using the SPAdes assembler version with the parameters --607 careful and a kmer range of "21, 29, 37, 45, 53, 61, 79, 87". The SPAdes assembler is optimized for 608 the assembly of relatively small eukaryotic genomes. We evaluated the completeness of the 609 assemblies using BUSCO v4.1.1 with the fungi odb10 gene test set (Simão et al., 2015). We finally 610 ran Quast v5.0.2 to retrieve assembly metrics including scaffolds of at least 1 kb (Mikheenko et al., 611 2018).

612

613 FUNGICIDE RESISTANCE ASSAY

To quantify susceptibility towards propiconazole we used a previously published microtiter plate assay dataset with 3 replicates performed for each isolate and concentration. Optical density was used to estimate growth rates under different fungicide concentrations (0, 0.00006, 0.00017, 0.0051, 0.0086, 0.015, 0.025, 0.042, 0.072, 0.20, 0.55, 1.5 mgL⁻¹) (Hartmann *et al.*, 2020). We calculated dose-response curves and estimated the half-maximal lethal concentration EC₅₀ with a 4-parameter logistics curve in the R package *drc* (Ritz & Streibig, 2005).

620

621 Data availability

622	Sequence data is deposited at the NCBI SRA under the accession numbers PRJNA327615,
623	PRJNA596434 and PRJNA178194. Transposable element consensus sequences are available from
624	https://github.com/crolllab/datasets.
625	
626	Author contributions
627	UO and DC conceived the study, UO, TW and DC designed analyses, UO, TB, TV and FEH
628	performed analyses, FEH, NKS, LNA, PK, CCM and BAM provided samples/datasets, BAM and
629	DC provided funding, UO and DC wrote the manuscript with input from co-authors. All authors
630	reviewed the manuscript and agreed on submission.
631	
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638	
639	Competing interests

- 640 We declare to have no competing interests.
- 641
- 642

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- 925

926 Figure Legends

927	
928	Figure 1: Robust discovery and validation of transposable element (TE) insertions: (A) General
929	analysis pipeline. (B) Read depth down-sampling analysis for one isolate per population with an
930	average coverage of the population. The vertical black line indicates the coverage at which on average
931	90% of the maximally detectable variants were recovered. Dashed black lines indicate the standard
932	error. The threshold for a minimal mean coverage was set at 15X (red line). (C) Validation of
933	insertions absent in the reference genome. (i) TE insertions that are not present in the reference
934	genome show a duplication of the target site and the part of the reads that covers the TE will not be
935	mapped against the reference genome. We thus expect reads to map to the TE surrounding region and
936	the target site duplication but not the TE itself. At the target site, a local duplication of read depth is
937	expected. (ii) We selected all reads in an interval of 100 bp up- and downstream including the target
938	site duplication to detect deviations in the number of reads terminating near the target site duplication.
939	(D) Validation of insertions present in the reference genome. (i) Analyses read coverage at target site
940	duplications. (ii) Decision map if a TE should be kept as a true insertion or rejected as a false positive.
941	Only predicted TE insertions that overlap evidence of split reads were kept as TE insertions in
942	downstream analyses. (E) Singleton validation using long-read PacBio sequencing. (i) Analysis if TE
943	insertions overlap with a detected insertion/deletion locus (Badet et al, 2021). (ii) Homology search of
944	the TE insertion flanking sequences based on the reference genome against PacBio reads. In addition,
945	the consensus sequence of the inserted TE was used for matches between the flanks.
946	Figure supplement 1. Validation of transposable element (TE) insertion predictions. (A) TEs not
947	present in the reference genome: distribution of additional TE hits found per locus after the outlier
948	test. Color indicates superfamilies. (B) TEs not present in the reference genome: distribution of
949	additional TE hits found per population after the outlier test. Colors indicate populations. (C) TEs
950	present in the reference genome: distribution of missing data per locus after the validation with
951	spliced junction reads. Missing data indicates that the TE was not predicted with ngs_te_mapper and
952	that there was no indication of spliced reads. The red line (=20 %) indicates the threshold for missing
953	data. TE loci with an amount of missing data > 20 % were completely excluded from further analyses.
954	Color indicates superfamily. (D) TEs present in the reference genome: detection of strong outlier
955	isolates with a high number of split reads. Color indicates the population.
956	Figure supplement 2. TE insertion validations for non-reference copies (Table).
957	Figure supplement 3. Establishment of transposable element (TE) loci with differing start and end
958	positions in the isolates. Distribution of length of distance for start position, end position and both
959	start and end combined after the correction.
960	Figure supplement 4. Bias for reads with a GC content lower than 30 % per population. Red lines

961 indicate the mean.

962 **Figure supplement 5.** TE consensus sequences (Table).

963

- 964 Figure 2: Transposable element (TE) landscape across populations. (A) Allele frequencies of the 965 TE insertions across all isolates. (B) TE insertions per Mb on core chromosomes (dark) and accessory 966 chromosomes (light). Dashed lines represent mean values. Blue: global mean of 75.65 insertions/Mb, 967 dark: core chromosome mean of 58 TEs/Mb, light: accessory chromosome mean of 102.24 968 insertions/Mb). (C) Number of TE insertions per family. (D) TE frequencies among isolates and copy 969 numbers across the genome. The blue line indicates the maximum number of isolates (n = 284). (E) 970 Allele frequency distribution of TE insertions into introns and exons. (F) Number of TE insertions 971 within 1 kb up- and downstream of genes on core chromosomes including introns and exons (100 bp 972 windows). The blue arrow indicates a gene schematic with exons and an intron, the green triangles 973 indicate TE insertions. The dotted blue line indicates no deviation from the expected value (i.e., mean 974 number of TEs per window). 975 Figure supplement 1. TEs in reference (Table). 976 **Figure supplement 2.** Validation of singleton insertions detected by mapped Illumina reads using 977 PacBio read alignments for confirmation. (A) Comparison of TE family copy numbers per isolate to 978 the number of copies found in the reference genome (IPO323). The color is indicating superfamilies. 979 This figure includes only TE families that were detected in any of the isolates used for validation. (B) 980 Confirmation of singleton TE insertions detected in the isolates CH99_SW5, CH99_SW39, 981 CH99_3D7, CH99_3D1, ISR92_Ar_4f, AUS01_1H8 and ORE90Ste_4A10 using aligned PacBio
 - 982 reads. Confirmed/not confirmed TE insertions are shown by TE family. (C) PacBio read coverage (in
 - 983 500 bp window) at singleton loci.
 - 984 Figure supplement 3. Presence absence matrix TE loci (Table).
 - 985 **Figure supplement 4.** Singletons (Table).

986 Figure supplement 5. TE insertion loci characteristics. (A) Number of TE insertions and density

- 987 (insertions per Mb) in accessory and core genes. (B) Allele frequencies of TEs genome-wide and
- 988 restricted to recombination hotspots. (C) TE insertion density and TE copy numbers within and
- 989 outside of recombination hotspots.
- 990 **Figure supplement 6.** Hierarchy superfamilies. (A) Number of transposable element (TE) insertions
- 991 per superfamily. Colors indicate the superfamily. (B) Number of TE loci and classification hierarchy.
- 992 (C) Comparison of mean genome sequencing coverage and the number of detected TEs with
- 993 ngs_te_mapper in isolates of the Middle East population. Dots indicate the coverage and colors
- 994 indicate the superfamily.
- 995

996 Figure 3: Differentiation in transposable element insertions frequencies across the genome. (A)

997 Global pairwise F_{ST} distributions shown across the 21 chromosomes. The red horizontal line indicates

- 998 the mean F_{ST} (= 0.0163). TEs with a strong local short-term frequency difference among populations
- are highlighted (blue: increase in Europe; green: increase in North America). (B) Allele frequency

- 1000 changes between the populations. The same TE loci as in panel A are highlighted. (C) Circos plot
- 1001 describing from the outside to the inside: The black line indicates chromosomal position in Mb. Blue
- 1002 bars indicate the gene density in windows of 100 kb with darker blue representing higher gene
- 1003 density. Red bars indicate the TE density in windows of 100 kb with a darker red representing higher
- 1004 TE density. Green triangles indicate positions of TE insertions with among population F_{ST} value
- 1005 shown on the y-axis.
- 1006 **Figure supplement 1.** Global pairwise FST distributions shown separately for the 21 chromosomes.
- 1007 The red horizontal line indicates the mean FST = 0.0163. Colors are according to the three main
- 1008 superfamilies (RLG, RLC, DHH).
- 1009
- 1010 Figure 4: Candidate adaptive transposable element (TE) insertions. (A) Distribution of all
- 1011 extremely differentiated TEs and their distance to the closest gene. Color indicates the superfamily.
- 1012 The stars indicate TE insertions not found in the reference genome. (B) Location of the RLG Luna
- 1013 TE insertion on chromosome 12 corresponding to its two closest genes. (C) Resistance against azole
- 1014 fungicides among isolates as a function of TE presence or absence. (D) Genomic niche of the
- 1015 RLG_Luna TE insertion on chromosome 12: F_{ST} values for each TE insertion, gene content (blue), TE
- 1016 content (green) and GC content (yellow). The grey section highlights the insertion site. (E) Number of
- 1017 RLG_Luna copies per isolate and population. (F) Frequency changes of RLG_Luna between the two
- 1018 North American populations compared to the other populations. Colors indicate the number of copies
- 1019 per chromosome. (G) Phylogenetic trees of the coding sequences of either the gene encoding the
- 1020 RTA1-like protein or the protein kinase domain. Isolates of the two North American populations and
- an additional 11 isolates from other populations not carrying the insertion are shown. Blue color
- 1022 indicates TE presence, yellow indicates TE absence.
- 1023 **Figure supplement 1.** Top loci information (Table).

1024 **Figure supplement 2.** Additional top loci. Six additional candidate adaptive transposable element

1025 (TE) insertions. Each row corresponds to a candidate, with the first five being candidates detected in

- 1026 the North American populations and the last one in the European populations. For each candidate, the
- 1027 direction of the TE and the direction, function and distance of the closest two genes are indicated. The
- 1028 middle column indicates the location of the TE in the genomic niche, with TE content, gene content
- 1029 and GC content for the surrounding windows. The third column indicates resistance levels towards
- 1030 azole antifungals for isolates with and without the TE insertion.
- 1031

1032 Figure 5: Population differentiation at transposable element (TE) and genome-wide SNP loci.

1033 (A) Sampling locations of the six populations. Middle East represents the region of origin of the

- 1034 pathogen. In North America, the two populations were collected at an interval of 25 years in the same
- 1035 field in Oregon. In Europe, two populations were collected at an interval of 17 years from two fields
- 1036 in Switzerland <20 km apart. Dark arrows indicate the historic colonization routes of the pathogen.

- 1037 (B) Principal component analysis (PCA) of 284 Zymoseptoria tritici isolates, based on 900,193
- 1038 genome-wide SNPs. (C) PCA of a reduced SNP data set with randomly selected 203 SNPs matching
- 1039 approximately the number of analyzed TE loci. (D) PCA based on 193 TE insertion loci. Loci with
- 1040 allele frequency < 5% are excluded.
- 1041 **Figure supplement 1.** Isolates (Table).
- 1042
- 1043 Figure 6: Global population structure of transposable element (TE) insertion polymorphism.
- 1044 (A) Total TE copies per isolate. Colors identify TE superfamilies. (B) TE copies per family and (C)
- 1045 superfamily. (D) TE insertion frequency spectrum per population. The curve fitting was performed
- 1046 with a self-starting Nls asymptomatic regression model (E). TE family copy numbers per isolate.
- 1047 Figure supplement 1. Population changes additional. Variation in transposable element (TE) content
- 1048 per isolate across populations. (A) Total TE copies per superfamily (colored) and per isolate only
- 1049 including LTR (long terminal repeat) TEs Copia and Gypsy. Color indicates the family. (B) Total TE
- 1050 copies per superfamily (colored) and per isolate only on the core chromosomes. (C) Total TE copies
- 1051 per superfamily (colored) and per isolate only on the accessory chromosomes.
- 1052 Figure supplement 2. Kolmogorof-Smirnov (Table).
- 1053 Figure supplement 3. Heatmap loci. (A) Presence (blue) and absence (yellow) matrix for all
- 1054 transposable element (TE) loci in all isolates per population. Colors on the left side indicate the
- superfamily. (B) Comparison of different genomic regions with and without TE insertions in IPO323.
- 1056

1057 Figure 7: Core genome size and transposable element (TE) evolution across populations. (A)

1058 BUSCO completeness variation among genome assemblies. Black lines indicate the mean genome

- 1059 size per population. (B) Genome-wide GC content variation. (C) Core genome size variation among
- 1060 the isolates of the populations (excluding accessory chromosomes). (D) Correlation of core genome
- 1061 size and number of detected TEs. (E) Correlation of core genome size and the cumulative length of all
- 1062 TEs detected as inserted. (F) Correlation of core genome size and genome-wide GC content. (G)
- 1063 Spearman correlation matrix of BUSCO completeness, core genome size, number of detected TEs and
- 1064 genome-wide GC content.
- 1065 Figure supplement 1. Genome size expansion. (A) Estimated length of TE insertions per isolate and
- 1066 population. (B) Genome size variation per population. (C) Percentage of TEs content variation
- 1067 compared to the variation in genome size. (D) TE contributions to genome size variation compared to
- 1068 full genome size.





















