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# A community-driven resource for genomic surveillance of Neisseria gonorrhoeae at Pathogenwatch — Source link $\square$

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#### A community-driven resource for genomic epidemiology and 1 antimicrobial resistance prediction of Neisseria gonorrhoeae 2 at Pathogenwatch 3

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45 Keywords: Neisseria gonorrhoeae, Pathogenwatch, public health, genomics, epidemiology, 46 surveillance, antimicrobial resistance.

## 48 Abstract

**Background:** Antimicrobial resistant (AMR) *Neisseria gonorrhoeae* is an urgent threat to public health, as strains resistant to at least one of the two last line antibiotics used in empiric therapy of gonorrhoea, ceftriaxone and azithromycin, have spread internationally. Whole genome sequencing (WGS) data can be used to identify new AMR clones, transmission networks and inform the development of point-of-care tests for antimicrobial susceptibility, novel antimicrobials and vaccines. Community driven tools that provide an easy access to and analysis of genomic and epidemiological data is the way forward for public health surveillance.

56 Methods: Here we present a public health focussed scheme for genomic epidemiology of N. 57 gonorrhoeae at Pathogenwatch (https://pathogen.watch/ngonorrhoeae). An international 58 advisory group of experts in epidemiology, public health, genetics and genomics of N. 59 gonorrhoeae was convened to inform on the utility of current and future analytics in the platform. 60 We implement backwards compatibility with MLST, NG-MAST and NG-STAR typing schemes as 61 well as an exhaustive library of genetic AMR determinants linked to a genotypic prediction of 62 resistance to eight antibiotics. A collection of over 12,000 N. gonorrhoeae genome sequences 63 from public archives has been quality-checked, assembled and made public together with 64 available metadata for contextualization.

65 **Results:** AMR prediction from genome data revealed specificity values over 99% for 66 azithromycin, ciprofloxacin and ceftriaxone and sensitivity values around 99% for benzylpenicillin 67 and tetracycline. A case study using the Pathogenwatch collection of *N. gonorrhoeae* public 68 genomes showed the global expansion of an azithromycin resistant lineage carrying a mosaic *mtr* 69 over at least the last 10 years, emphasizing the power of Pathogenwatch to explore and evaluate 70 genomic epidemiology questions of public health concern.

Conclusions: The *N. gonorrhoeae* scheme in Pathogenwatch provides customized bioinformatic pipelines guided by expert opinion that can be adapted to public health agencies and departments with little expertise in bioinformatics and lower resourced settings with internet connection but limited computational infrastructure. The advisory group will assess and identify ongoing public health needs in the field of gonorrhoea, particularly regarding gonococcal AMR, in order to further enhance utility with modified or new analytic methods.

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#### 82 Background

83 Antimicrobial resistance (AMR) is an urgent threat to public health. Neisseria gonorrhoeae, the 84 strictly human pathogen causing the sexually-transmitted infection (STI) gonorrhoea, has 85 developed or acquired resistance to the last-line antibiotics used in empiric therapy to treat the 86 infection, and thus has become one of the major global priorities in order to tackle AMR. In 2017, 87 due to the increase in AMR infections and the absence of an effective vaccine, the World Health 88 Organization (WHO) included N. gonorrhoeae as a high priority pathogen in need of research and 89 development of new antimicrobials and ideally a vaccine (1). In 2019, the Centers for Disease 90 Control and Prevention (CDC) again included the gonococcus on the list of urgent threats in the 91 United States (2). The most recent WHO estimates from 2016 indicate an annual global incidence 92 of 87 million cases of gonorrhoea among adults (3, 4). Untreated cases can develop complications 93 including an increased acquisition and transmission of HIV. In women, long-term infections can 94 cause infertility, pelvic inflammatory disease, ectopic pregnancy, miscarriage or premature labour 95 (5). Infections during pregnancy can transmit to newborns at birth causing eye damage that can 96 have permanent effects on vision (6).

97 Strains of *N. gonorrhoeae* resistant to every recommended treatment have rapidly emerged, 98 including resistance to penicillins, tetracyclines, fluoroquinolones, macrolides and the extended-99 spectrum cephalosporins (ESCs) (5-8). The current recommended treatment in many countries 100 is a dual therapy with injectable ceftriaxone plus oral azithromycin, although reports of decreased 101 susceptibility to ceftriaxone as well as azithromycin resistance have increased globally (7, 8). One 102 case of failure of dual treatment was reported in 2016 in the United Kingdom (UK) (9). Additionally, 103 in 2018 a gonococcal strain with resistance to ceftriaxone combined with high-level resistance to 104 azithromycin was detected in both the UK and Australia (10). The transmission of a ceftriaxone-105 resistant clone (FC428) has been documented internationally since 2015, raising concerns about 106 the long-term effectiveness of the current treatment in the absence of an available alternative 107 (11). In some countries such as in Japan, China and since 2019 in the UK, a single dose of 108 ceftriaxone 1 gram is the recommended treatment due to the increasing incidence of azithromycin 109 resistance in N. gonorrhoeae and other STI pathogens such as Mycoplasma genitalium (12). 110 Extensive investigations have been ongoing for years to unveil the genetic mechanisms that explain most of the observed susceptibility patterns for the main classes of antimicrobials for N. 111 gonorrhoeae. For ciprofloxacin, nearly all resistant strains have the GyrA S91F amino acid 112 113 alteration (13-15), however, resistance prediction from genomic data is not as straightforward for other antibiotics. Known resistance mechanisms often involve additive or suppressive effects as 114 115 well as epistatic interactions that all together explain just part of the observed phenotypic 116 resistance. For example, there is good evidence that many mosaic structures of the penA gene 117 are associated with decreased susceptibility to ESCs (16, 17), however, mosaics do not explain 118 all cases of ESC resistance, especially for ceftriaxone, and some mosaic penA alleles do not cause decreased susceptibility or resistance to this antibiotic (16-19). On top of these, variants 119 120 that overexpress the MtrCDE efflux pump, mutations in *porB1b* that reduce drug influx and non-121 mosaic mutations in penicillin-binding proteins also contribute to decreased susceptibility to ESCs 122 (20). Furthermore, mutations in the rpoB and rpoD genes, encoding subunits of the RNA 123 polymerase, have been recently related to resistance to ESCs in clinical N. gonorrhoeae isolates 124 (21). Mutations in the 23S rRNA gene (A2045G and C2597T in N. gonorrhoeae nomenclature, 125 coordinates from the WHO 2016 reference panel (22), A2059G and C2611T in Escherichia coli) 126 are frequently associated with azithromycin resistance, as do variants in *mtrR* or its promoter that 127 increase the expression of the MtrCDE efflux pump (5). Recently, epistatic interactions between 128 a mosaic *mtr* promoter region and a mosaic *mtrD* gene have also been reported to increase the 129 expression of this pump, contributing to macrolide resistance (23, 24). Mutations in rpID have also 130 been associated with reduced susceptibility to this antibiotic (25) and contrarily, loss-of-function 131 mutations in *mtrC* have been linked to increased susceptibility to several antibiotics including 132 azithromycin (26). Thus, we can relatively confidently predict decreased susceptibility or 133 resistance to an antimicrobial using the current known genetic mechanisms, however, phenotypic 134 testing is still necessary to detect resistant cases caused by unknown or novel mechanisms. 135 These inconsistencies with the genomic data will allow the discovery of these new mechanisms, 136 which will keep improving the resistance predictions from WGS.

137 A myriad of methods have been used to discriminate among strains of N. gonorrhoeae, from 138 phenotypic to DNA-based techniques (27), but whole genome sequencing (WGS) can provide 139 the complete genome information of a bacterial strain. The cost of amplifying all loci of the different 140 typing schemes via nucleic acid amplification and traditional Sanger sequencing can be more 141 expensive than the cost of WGS of one bacterial genome in many settings. With WGS, multiple 142 genetic AMR mechanisms as well as virulence and typing regions can be targeted simultaneously with the appropriate bioinformatic tools and pipelines. It also provides a significant improvement 143 144 in resolution and accuracy over traditional molecular epidemiology and typing methods, allowing a genome-wide comparison of strains that can: identify AMR clones, outbreaks, transmission 145 146 networks, national and international spread, known and novel resistance mechanisms as well as 147 also inform on the development of point-of-care tests for antimicrobial susceptibility, novel 148 antimicrobials and vaccines (28, 29). However, implementation of WGS for genomic surveillance 149 poses practical challenges, especially for Low- and Middle-Income Countries (LMICs), due to the 150 need of a major investment to acquire and maintain the required infrastructure.

151 WGS produces a very high volume of data that needs to be pre-processed and analysed using 152 bioinformatics. Bioinformatics expertise is not always readily available in laboratory and public 153 health settings, and currently there are no international standards and proficiency trials for which 154 algorithms to use to process WGS data. There are several open-source tools specialised in each 155 step of the pipeline as well as proprietary software containing workflows that simplify the analyses. 156 However, these are less customizable and may not be affordable for all (30, 31). Choosing the 157 best algorithms and parameters when analysing genomic data is not straightforward as it requires 158 a fair knowledge of the pathogen under study and its genome diversity. Multiple databases 159 containing genetic determinants of AMR for bacterial pathogens are available (30, 31), however, 160 choosing which one is most complete for a particular organism frequently requires an extensive 161 literature search. Public access web-based species-specific tools and AMR databases revised 162 and curated by experts would be the most approachable option for both well-resourced and LMICs 163 with a reliable internet connection. Very importantly though, the full benefits of using WGS for

both molecular epidemiology and AMR prediction can only be achieved if the WGS data are linked
to phenotypic data for the gonococcal isolates and, as much as feasible, clinical and
epidemiological data for the patients.

167 Here, we present a public health focussed system to facilitate genomic epidemiology of N. 168 gonorrhoeae within Pathogenwatch (https://pathogen.watch/ngonorrhoeae), which includes the 169 latest analytics for typing, detection of genetic AMR determinants and prediction of AMR from N. 170 gonorrhoeae genome data, linked to metadata where available, as well as a collection of over 171 12,000 gonococcal genomes from public archives for contextualization. We formed an advisory 172 group including experts in the field of *N. gonorrhoeae* epidemiology, public health, AMR, genetics 173 and genomics to consult on the development and design of the tool, such as the analytics and 174 genetic AMR mechanisms to include, in order to adapt the platform for ongoing public health 175 needs. We present this scheme as a community-steered model for genomic surveillance that can 176 be applied to other pathogens.

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# 178 Methods

#### 179 The Pathogenwatch platform: technical summary

180 Pathogenwatch is a web-based platform with several different components. The main interface is 181 a React (32) single-page application with a style based on Material Design Lite (33). Phylogenetic 182 trees are plotted using Phylocanvas (34), maps using Leaflet (35) and networks with Sigma (36). 183 The back end is written in Node is and contains an API service for the user interface and four 184 "Runner" services for the following analyses: species prediction, single-genome analyses, tree building and core genome multi-locus sequence typing (cgMLST) clustering. Docker containers 185 are used for queuing tasks, streaming input or result files through standard input and storing 186 187 JSON data from standard output. A MongoDB cluster is used for data storage and task 188 queuing/synchronisation. Pathogenwatch shares some visualization components with Microreact

(37), such as those associated with the phylogenetic tree and the map. However, Pathogenwatchincludes an analytical framework which is unique to this platform.

#### 191 Generation of the N. gonorrhoeae core genome library

192 Pathogenwatch implements a library of core genome sequences for several supported organisms. 193 In the case of *N. gonorrhoeae*, a core gene set was built from the 14 finished reference genomes 194 that constitute the 2016 WHO reference strain panel (22) using the pangenome analysis tool 195 Roary (38) as described in Harris et al (2018) (15). Briefly, the minimum percentage of identity for 196 blastp was set to 97% and the resulting core genes were aligned individually using MAFFT. The 197 resulting genes with a percentage of identity above 99% were post-processed as described in 198 (39). Representatives for each family were selected by choosing the sequence with the fewest 199 differences to the others on average and searched using tblastn (percentage of identity  $\geq 80\%$ , 200 E-value <= 1e-35) against the 14 high quality reference genomes. Families without a complete match in every reference (100% coverage) or had multiple matches were removed. Overlapping 201 202 genes from each reference were merged into pseudocontigs and grouped by gene composition. 203 For each family, a representative was selected as before and searched/filtered using the 204 references as before. The final core gene set contains 1,542 sequences that span a total of 205 1,470,119 nucleotides (approximately 67% of a typical *N. gonorrhoeae* genome length, 2.2Mb). 206 A BLAST database was constructed from these core segments and used to profile new 207 assemblies.

## 208 **Profiling new assemblies**

New genome assemblies can be uploaded by a user (drag and drop) or calculated from highthroughput short read data directly within Pathogenwatch using SPAdes (40) as described in (41).

A taxonomy assignment step for species identification is performed on the uploaded assemblies by using Speciator (42). New assemblies are then queried against a species-specific BLAST database using blastn. For *N. gonorrhoeae*, every core loci needs to match at least 80% of its length to be considered as present. Further filtering steps are applied to remove loci that can be problematic for tree building, such as paralogs or loci with unusually large number of variant sites compared to an estimated substitution rate on the rest of the genome, as described in (43). The overall substitution rate is calculated as the number of total differences in the core library divided by the total number of nucleotides. Indels are ignored to minimise the noise that could be caused by assembly or sequencing errors. The expected number of substitutions per locus is determined by multiplying this substitution rate by the length of the representative sequence.

The number of substitutions observed for each locus between the new assembly and the reference sequence are scaled to the total number of nucleotides that match the core library, creating a pairwise score that is saved on a distance matrix and is used for Neighbour-Joining tree construction, as described in (44).

## 225 Algorithms for sequence typing and cgMLST clustering

226 Alleles and sequence types (STs) for Multi-Locus Sequence Typing (MLST) (45) and cgMLST 227 (core genome MLST, N. gonorrhoeae cgMLST v1.0) (46) were obtained from PubMLST (47, 48), 228 for N. gonorrhoeae Multi-Antigen Sequence Typing (NG-MAST) (49) from (50) and for N. 229 gonorrhoeae Sequence Typing for Antimicrobial Resistance (NG-STAR) (51) from (52) (Table 1). 230 A search tool implemented as part of Pathogenwatch is used to make the assignments for MLST, 231 cgMLST and NG-STAR, while NGMASTER (53) is used for NG-MAST. Briefly, exact matches to 232 known alleles are searched for, while novel sequences are assigned a unique identifier. The 233 combination of alleles is used to assign a ST as described in (54). Databases are regularly 234 updated and novel alleles and STs should be submitted by the user to the corresponding schemes 235 for designation.

cgMLST typing information is used for clustering individual genomes with others in the
Pathogenwatch database using single linkage clustering as described in (55). Users can select
the clustering threshold (i.e. number of loci with differing alleles) and a network graph based on
the SLINK (56) algorithm is calculated within individual genome reports.

#### 241 AMR library and detection of genetic AMR determinants

242 Genes and point mutations (single nucleotide polymorphisms (SNPs) and indels) were detected 243 using Pathogenwatch AMR v2.4.9 (57). Pathogenwatch AMR also provides a prediction of AMR 244 phenotype inferred from the combination of identified mechanisms. Genetic determinants 245 described in the literature as involved in AMR in N. gonorrhoeae were collated into a library in 246 TOML format (version 0.0.11). A test dataset containing 3,987 isolates from 13 studies (15, 18, 247 22, 58-67) (Additional file 1: Table S1) providing minimum inhibitory concentration (MIC) 248 information for six antibiotics (benzylpenicillin, tetracycline, ciprofloxacin, cefixime, ceftriaxone 249 and azithromycin) was used to benchmark and to curate this library. A validation benchmark was 250 posteriorly run with a dataset of 1,607 isolates from 3 other publications (68-70) with MIC 251 information for the same six antibiotics plus spectinomycin (Additional file 1: Table S1). EUCAST 252 clinical breakpoints v9.0 (71) were used to define susceptibility (S), susceptibility with an 253 increased exposure (I) or resistance (R) (SIR) categorical interpretations of MICs for all antibiotics 254 except for azithromycin, for which the EUCAST epidemiological cut-off (ECOFF) was used to 255 define non-susceptibility/resistance (ECOFF>1mg/L). As a result of the benchmark analyses, 256 sensitivity, specificity and positive/negative predictive values (PPV/NPV) were obtained for the 257 AMR mechanisms implemented in the library and, globally, for each of the antibiotics. Confidence 258 intervals (95%) for these statistics were calculated using the epi.tests function in the epiR R 259 package v1.0-14 (72). Individual or combined AMR mechanisms with a PPV below 15% were 260 discarded from the library to optimise the overall predictive values. Visual representations of the 261 observed ranges of MIC values for a particular antibiotic for each of the observed combinations 262 of genetic AMR mechanisms on the test dataset were used to identify and assess combinations 263 of mechanisms that have an additive or suppressive effect on AMR. These were included in the 264 library.

As part of the accuracy testing of the AMR library, we ran the 2016 WHO *N. gonorrhoeae* reference genomes 2016 panel (n=14) through Pathogenwatch and compared the detected list of genetic AMR mechanisms with the list published in the original study (22). For the WHO U

strain, a discrepancy on a mutation in *parC* was further investigated by mapping the original raw Illumina data (European Nucleotide Archive (ENA) run accession ERR449479) to the reference genome assembly (ENA genome accession LT592159.1) and visualized using Artemis (73).

In short-read assemblies, the four copies of the 23S rRNA gene are collapsed into one, thus the detection of the A2045G and C2597T mutations is dependent on the consensus bases resulting from the number of mutated copies (63, 66, 74).

#### 274 Quality check and assembly of public sequencing data

275 Public *N. gonorrhoeae* genomes with geolocation data were obtained from the ENA in November 276 2019. This list was complemented by an exhaustive literature search of studies on N. gonorrhoeae 277 genomics without metadata submitted to the ENA but instead made available as supplementary 278 information in the corresponding publications. Raw paired-end short read data from a list of 279 12,192 isolates was processed with the GHRU assembly pipeline v1.5.4 (75). This pipeline runs 280 a Nextflow workflow to quality-check (QC) paired-end short read fastq files before and after 281 filtering and trimming, assembles the data and quality-checks the resulting assembly. Results 282 from the pipeline are provided in Additional file 2. In this pipeline, QC of short reads was performed 283 using FastQC v0.11.8 (76). Trimming was done with Trimmomatic v0.38 (77) by cutting bases 284 from the start and end of reads if they were below a Phred score of 25, trimming using a sliding 285 window of size 4 and cutting once the average quality within the window fell below a Phred score 286 of 20. Only reads with length above a third of the original minimum read length were kept for 287 further analyses. After trimming, reads were corrected using the kmer-based approach 288 implemented in Lighter v1.1.1 (78) with a kmer length of 32 bp and a maximum number of 289 corrections allowed within a 20 bp window of 1. ConFindr v0.7.2 was used to assess intra- and 290 inter-species contamination (79). Mash v2.1 (80) was applied to estimate genome size using a 291 kmer size of 32 bp and Seqtk v1.3 (81) to down sample fastq files if the depth of coverage was 292 above 100x. Flash v1.2.11 (82) was used to merge reads with a minimum overlap length of 20 bp 293 and a maximum overlap of 100 bp to facilitate the subsequent assembly process. SPAdes v3.12 (40) was used for genome assembly with the --careful option selected to reduce the number of
mismatches and short indels with a range of kmer lengths depending on the minimum read length.
The final assemblies were quality-checked using Quast v5.0.2 (83) and ran through the species
identification tool Bactinspector (84). QC conditions were assessed and summarised using
Qualifyr (85).

Fastq files with poor quality in which the trimming and filtering step discarded all reads from either one or both pairs were excluded from the analyses because the assembly pipeline is optimised for paired-end data. Assemblies with an N50 below 25,000 bp, a number of contigs above 300, a total assembly length above 2.5 Mb or a percentage of contamination above 5% were also excluded.

#### 304 Metadata for public genomes

305 Geolocation data (mainly country), collection dates (day, month and year when available), ENA 306 project accession and associated Pubmed ID were obtained from the ENA API for all the genomes 307 in the pipeline (86). A manual extensive literature search was performed to identify the 308 publications containing the selected genomes. In order to complete published studies as much 309 as possible, extra genomes were downloaded and added to the dataset. Metadata for the final 310 set was completed with the information contained in supplementary tables on the corresponding 311 publications, including phenotypic antimicrobial susceptibility data. Submission date was 312 considered instead of collection date when the latter was not available, however, this occurred in 313 only a few cases (<0.5%).

#### 314 Creation of the N. gonorrhoeae Pathogenwatch Scientific Steering Group

International experts in the field of *N. gonorrhoeae* AMR, microbiology, genetics, genomics, epidemiology and public health were approached and agreed to participate as members of the *N. gonorrhoeae* Pathogenwatch Scientific Steering Group' in order to discuss the analytics in Pathogenwatch and make sure they met the current needs of the public health and scientific community. During the updates made to the platform and the preparation of this manuscript, these experts participated in virtual sessions to discuss the list of genetic AMR determinants and their association with SIR categories (Table 2) based on experimental and/or computational evidence. Some of the members of the group had previously been directly involved in many of these studies. Other current and future updates were also discussed, such as the inclusion of the NG-STAR typing scheme (51) and the organization of published genomes into public collections, data sharing, privacy and the interconnectivity of Pathogenwatch with other platforms, such as PubMLST (48) or the ENA. The group will regularly discuss new updates to the platform.

#### 327 Data sharing and privacy

328 Sequencing data and metadata files uploaded to Pathogenwatch by the user are kept within the 329 user's private account. Genomes can be grouped into collections and these can be toggled 330 between private and accessible to collaborators via a URL. Collection URLs include a 12-letter 331 random string to secure them against brute force searching. Setting a collection to 'off-line mode' 332 allows users to work in challenging network conditions, which may be beneficial in LMICs - all 333 data are held within the browser. Users can also integrate private and potentially confidential 334 metadata into the display without uploading it to the Pathogenwatch servers (locally within the 335 browser on a user's machine).

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#### 337 Results

#### 338 N. gonorrhoeae genome analytics in Pathogenwatch

Pathogenwatch is a web-based platform for epidemiological surveillance using genome sequencing data. After upload, different analytics are run simultaneously (Figure 1): cgMLST (46), MLST (45), NG-MAST (49) and NG-STAR (51) typing schemes (Table 1), a genotypic prediction of phenotypic resistance using a customized AMR library (Table 2) that includes known genetic AMR mechanisms for 8 antimicrobials, as well as statistics on the quality of the assemblies (Additional file 3: Figure S1). These analytical features differentiate Pathogenwatch from a parallel

345 platform from the same group, Microreact (37), which shares one of the main layouts with 346 Pathogenwatch (a phylogenetic tree, a map and a table or timeline), but it is intended for 347 visualization of pre-computed phylogenetic trees with accompanying metadata, while 348 Pathogenwatch also includes analytical tools.

349 Genomes from one or multiple studies can be grouped into collections (Figure 2 and Additional 350 file 3: Figure S2), and the genomic data are automatically processed by comparing with a core N. 351 gonorrhoeae genome built from WHO reference strain genomes (15, 22). A phylogenetic tree is 352 obtained as a result, representing the genetic relationship among the isolates in the collection. 353 Metadata can be uploaded at the same time as the genome data, and if the collection location 354 coordinates for an isolate are provided, this information is plotted into a map (Additional file 3: 355 Figure S1). If date or year of isolation is also provided, this information is represented in a timeline. 356 The three panels on the main collection layout - the tree, the map and a table or timeline – are 357 functionally integrated so filters and selections made by the user update all of them 358 simultaneously. Users can also easily switch among the metadata and the results of the main 359 analytics: typing, genome assembly statistics, genotypic AMR prediction, AMR-associated SNPs, 360 AMR-associated genes and the timeline (Additional file 3: Figure S1). cgMLST is used for finding 361 close genomes in the database based on allele differences to one individual isolate (Additional 362 file 3: Figure S3). A video demonstrating the usage and main features of Pathogenwatch is 363 available (87). Notes on data sharing and privacy are available in the Methods section.

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Figure 1. Main workflow in Pathogenwatch. New genomes can be uploaded and combined with public data for contextualisation. The collection view allows data exploration through a combined phylogenetic tree, a map, a timeline and the metadata table, which can be switched to show typing information (Multi-Locus Sequence Typing, MLST; *N. gonorrhoeae* Sequence Typing for Antimicrobial Resistance, NG-STAR; and *N. gonorrhoeae* Multi-Antigen Sequence Typing, NG-MAST) as well as known genetic AMR mechanisms for eight antibiotics. Genome reports summarise the metadata, typing and AMR marker results for individual isolates and allow finding other close genomes in Pathogenwatch based on core genome MLST (cgMLST). SNPs: single nucleotide polymorphisms.



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Figure 2. Main display of a Pathogenwatch collection, showing a phylogenetic tree, a map and a table of SNPs associated with AMR of 395 *N. gonorrhoeae* genomes from a global study (64, 88). Isolates carrying three mosaic *penA* marker mutations are marked in red in the tree and the map. The table can be switched to show the metadata, a timeline, typing results (Multi-Locus Sequence Typing, MLST; *N. gonorrhoeae* Sequence Typing for Antimicrobial Resistance, NG-STAR and *N. gonorrhoeae* Multi-Antigen Sequence Typing, NG-MAST) as well as AMR analytics (known genetic mechanisms and genotypic AMR prediction) implemented in the platform. Further detail is shown in Additional file 3: Figure S1.

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398 Table 1. *N. gonorrhoeae* sequence typing schemes implemented in Pathogenwatch.

Typing scheme*	Loci (number)	Note	Pathogenwatch implementation	References
cgMLST	(N=1,649)	<i>N. gonorrhoeae</i> cgMLST v1.0	Typing algorithm, database from PubMLST	(46-48, 89)
MLST	abcZ, adk, aroE, fumC, gdh, pdhC, pgm (N=7)	Housekeeping genes in <i>Neisseria</i> spp.	In-house typing tool, database from PubMLST	(45, 47, 48, 89)
NG-MAST	porB, tbpB (N=2)	Genes encoding highly-variable membrane proteins	NG-MASTER, database from NG-MAST website	(49, 50, 53)
NG-STAR	penA, mtrR, porB, ponA, gyrA, parC, 23S rDNA (N=7)	Genes involved in antimicrobial resistance	In-house typing tool, database from NG-STAR website	(51, 52, 89)

400 \* Typing scheme: cgMLST = core genome Multi-Locus Sequence Typing, MLST = Multi-Locus Sequence Typing, NG-

401 MAST = *N. gonorrhoeae* Multi-Antigen Sequence Typing, NG-STAR = *N. gonorrhoeae* Sequence Typing for 402 Antimicrobial Resistance.

Table 2. List of *N. gonorrhoeae* genetic antimicrobial resistance (AMR) determinants in Pathogenwatch. References that report evidence of association of each mechanism to AMR in clinical isolates and/or where their role on AMR has been confirmed in the laboratory through, e.g. transformation experiments, are included in the table. Effect: R = resistance, I = susceptibility but increased exposure, A = additive effect, N = negative effect. R and I follow the EUCAST clinical breakpoints except for azithromycin, for which the epidemiological cut-off (ECOFF) is reported and used instead.

Antibiotic (MIC breakpoint mg/L)	Genetic AMR determinants	Effect	Evidence (References)
Azithromycin (R: MIC>1, ECOFF)	23S rDNA 2045A>G substitution (2059A>G in <i>E. coli</i> ) 23S rDNA 2597C>T substitution (2611C>T in <i>E. coli</i> ) ermA, ermB, ermC, ermF genes ereA, ereB genes mefA gene macAB promoter -48G>T substitution* mtrR promoter mosaic**	R R R R R R	(74) (90) (91, 92) (22) (92, 93) (94)
	<i>N. meningitidis</i> -like mosaic (n=1) <i>N. lactamica</i> -like mosaic (n=2) <i>mtrD</i> mosaic**	R R	(23) (23)
	N. meningitidis-like mosaic (n=1) N. lactamica-like mosaic (n=2) mtrR promoter -57delA* mtrR G45D mtrC loss-of-function rpIV ARAK tandem duplication (position 90) rpIV KGPSLK tandem duplication (position 83) rpID G70D	R A A N R A	(23) (23) (95, 96) (97, 98) (26) (18) (18) (25)
Ceftriaxone*** (R: MIC>0.125)	<i>penA</i> mosaic (A311V, I312M, V316P/T, T483S and G545S) <i>penA</i> V316P, T483S, A501P/V, G542S <i>rpoB</i> P157L, G158V, R201H <i>rpoD</i> D92-95 deletion, E98K	R R R I	(99-101) (99, 100) (21) (21)
Cefixime*** (R: MIC>0.125)	<i>mtrR</i> G45D <i>penA</i> mosaic (I312M, V316T, G545S) <i>penA</i> mosaic (A311V, I312M, V316P/T, T483S and G545S) <i>penA</i> V316P, T483S, A501P <i>rpoB</i> P157L, G158V, R201H <i>rpoD</i> D92-95 deletion, E98K	A R I I I	(97, 98) (99-101) (99-101) (99, 100) (21) (21)
Ciprofloxacin (I: 0.03 <mic≤0.06; R: MIC&gt;0.06)</mic≤0.06; 	gyrA S91F, D95A/N gyrA D95G norM promoter -7A>G, -104C>T substitutions* parC D86N, S87R parC S87I/N, S88P, E91K parE G410V	R I I R I I	(102) (102) (103) (102) (102) (104)
Tetracycline**** (I: 0.5 <mic≤1; R: MIC&gt;1)</mic≤1; 	<i>mtrR</i> A39T, G45D <i>mtrR</i> loss-of-function <i>mtrR</i> promoter -56A>C substitution, -57delA deletion* <i>mtrR</i> promoter -131G>A ( <i>mtrC</i> -120G>A substitution, <i>mtr120</i> )* <i>rpsJ</i> V57M <i>tetM</i> gene	A I I I R	(97, 98) (22) (23, 95, 96) (97) (105) (106)

Penicillins (I: 0.06 <mic≤1; R: MIC&gt;1) Spectinomycin</mic≤1; 	<i>blaTEM</i> gene <i>mtrR</i> G45D <i>mtrR</i> A39T <i>mtrR</i> loss-of-function <i>mtrR</i> promoter -56A>C, -57deIA* <i>mtrR</i> promoter -131G>A ( <i>mtrC</i> -120G>A substitution, <i>mtr120</i> )* <i>penA</i> I312M, V316P/T, ins346D, T483S, A501P/T/V, G542S, G545S, P551S <i>penA</i> mosaic (I312M, V316T, G545S) <i>ponA1</i> L421P <i>porB1b</i> G120K, A121N/D 16S rDNA 1184C>T (1192C>T in <i>E. coli</i> )	R I I I I I R	(107) (97, 98) (97) (22) (23, 96) (97) (99, 100) (99-101) (108) (109) (110)
(R: MIC>64)	rpsE T24P rpsE V27- deletion, K28E	R R/A	(111) (111)
Sulfonamides	folP R228S	R	(22, 112)

# 409

410 \*Nomenclature of the mutations on the macAB, mtrR and norM promoter regions is based on N. gonorrhoeae coordinates considering 411 the distance from the start of the macAB, mtrR and norM genes, respectively. \*\*Note that mosaics are caused by recombination events, 412 which can have variable breakpoints with different effects on azithromycin MIC if any. In this version, we have included the three 413 mosaics described by Wadsworth et al. (23), but the list will be expanded as new mosaic mtr (intergenic region between mtrR and 414 mtrC) and mtrD alleles having an effect on azithromycin MICs are published. \*\*\*The list of genetic AMR mechanisms for the ESCs 415 ceftriaxone and cefixime do not include all known porB1b or mtrR-associated variants as their effect was found not to be relevant in 416 increasing MIC on the benchmark analyses for phenotypic AMR prediction purposes despite the experimental evidence reported in 417 Zhao et al. (113). In case of strains carrying penA-associated mutations, their immediate predicted phenotype is that of those carrying 418 penA-associated variants. \*\*\*\*The list of genetic AMR mechanisms for tetracycline does not include porB1b mutations as their effect 419 was found not to be relevant in increasing MIC on the benchmark analyses for phenotypic AMR prediction purposes. \*\*\*\*\*Sulfonamides 420 are not a treatment alternative for gonorrhoea, however the foIP R228S mutation is kept in this version of the library for surveillance 421 purposes.

## 422 Library of genetic AMR mechanisms: genotypic and phenotypic benchmarks

423 We compiled described genetic AMR mechanisms previously reported for N. gonorrhoeae up to 424 the writing of this manuscript into the AMR library in Pathogenwatch (Table 2). A genotypic 425 accuracy testing of the AMR library was performed using the 14 N. gonorrhoeae reference 426 genomes from the WHO 2016 panel (22), which were uploaded into Pathogenwatch. All the 427 genetic AMR determinants described as present in these isolates and implemented in the 428 Pathogenwatch AMR library were obtained as a result (Additional file 1: Table S2). Only one 429 discrepancy was found when compared to the original publication. The WHO U strain was 430 reported as carrying a parC S87W mutation. However, mapping the original Illumina data from 431 this isolate with the final genome assembly revealed that this strain carries a wild type allele 432 (Additional file 3: Figure S4). MLST and NG-MAST types were the same as those reported in the 433 original publication (note that NG-STAR was not available at that time) and the porA mutant gene 434 was found in WHO U as previously described. This mutant porA has nearly a 95% nucleotide

identity to *N. meningitidis* and 89% to *N. gonorrhoeae*, and it is included as screening because it
has previously been shown to cause false negative results in some molecular detection tests for *N. gonorrhoeae* (114).

438 Then, we also performed a genotypic-phenotypic benchmark using a test dataset of 3,987 N. 439 gonorrhoeae isolates from 13 different studies containing MIC information for at least part of the 440 following six antibiotics: ceftriaxone, cefixime, azithromycin, ciprofloxacin, benzylpenicillin and 441 tetracycline (Additional file 1: Table S1). EUCAST clinical breakpoints were applied for five of the 442 antimicrobials except for azithromycin, for which the adoption of an ECOFF>1 mg/L is now 443 recommended to distinguish isolates with azithromycin resistance determinants, instead of a 444 clinical resistance breakpoint (115, 116). A visualization of the range of MICs on each particular 445 combination of genetic AMR mechanisms observed on the isolates from the benchmark test 446 dataset (Figure 3a-b and Additional file 3: Figures S5-S10) revealed combinations that show an 447 additive effect on AMR. These combinations were included in the AMR library to improve the 448 accuracy of the genotypic prediction. For example, rpsJ V57M and some mtrR-associated 449 mutations individually are associated with a decreased susceptibility or intermediate resistance 450 to tetracycline (MICs of 0.5-1 mg/L), however, a combination of these variants can increase MICs 451 above the EUCAST resistance breakpoint for tetracycline (MICs>1 mg/L) (Additional file 3: Figure 452 S9). This is the case of the combination of *rpsJ* V57M with the *mtrR* promoter -57delA mutation 453 (N=681 isolates, 94.9% positive predictive value, PPV) or with mtrR promoter -57delA and mtrR 454 G45D (N=83 isolates, 93.9% PPV). Several combinations of penA, ponA1, mtrR and porB1b 455 mutations were observed to be able to increase the benzylpenicillin MIC above the resistant 456 threshold in most of the cases (Additional file 3: Figure S10). This is the case of the porB1b 457 mutations combined with mtrR A39T (N=31 isolates, 100% PPV), with the mtrR promoter -57delA 458 deletion (N=286 isolates, 96.5% PPV) or with *mtrR* promoter -57delA and *ponA1* L421P (N=269 459 isolates, 96.3%). Despite mosaic penA not being a main driver of resistance to penicillins, a 460 combination of the *porB1b* mutations with the three main mosaic *penA* mutations (G545S, I312M 461 and V316T) was also associated with a resistant phenotype in all cases (N=17 isolates, 100%

462 PPV). A recent publication showed that loss-of-function mutations in *mtrC* increased susceptibility 463 to azithromycin and are associated with isolates from the cervical environment (26). We included 464 the presence of a disrupted mtrC as a modifier of antimicrobial susceptibility in the presence of 465 an mtr mosaic, as we did not have enough evidence from the test dataset to assess the MIC 466 ranges of isolates with the 23S rDNA A2045G and C2597T mutations with and without a disrupted 467 mtrC gene.





470 Figure 3. Distribution of minimum inhibitory concentration (MIC) values (mg/L) for the last-line antibiotics for N. 471 gonorrhoeae azithromycin (a) and ceftriaxone (b) in a collection of 3,987 N. gonorrhoeae isolates with different 472 combinations of genetic antimicrobial resistance (AMR) mechanisms. Only combinations observed in at least 5 isolates 473 are shown (see Additional file 3: Figure S5-S10 for expanded plots for six antibiotics). Dashed horizontal lines on the 474 violin plots mark the EUCAST epidemiological cut-off (ECOFF) for azithromycin and EUCAST clinical breakpoint for 475 ceftriaxone. Point colours inside violins represent the genotypic AMR prediction by Pathogenwatch on each 476 combination of mechanisms (indicated by black circles connected vertically; horizontal thick grey lines connect 477 combinations of mechanisms that share an individual determinant). Barplots on the top show the abundance of isolates 478 with each combination of mechanisms. Bar colours represent the differences between the predicted and the observed 479 SIR (i.e. red for a predicted susceptible mechanism when the observed phenotype is resistant). (c) Radar plots 480 comparing the sensitivity, specificity, positive and negative predictive values (PPV/NPV) for six antibiotics for the test

481 and validation benchmark analyses. AZM = Azithromycin, CFM = Cefixime, CIP = Ciprofloxacin, CRO = Ceftriaxone,
 482 PEN = Benzylpenicillin, TET = Tetracycline.

483

Results from the benchmark (Additional file 1: Table S3) show sensitivity values (true positive 484 485 rates. TP/(TP+FN): TP=True Positives. FN=False Negatives) above 96% for tetracycline (99.2%). 486 benzylpenicillin (98.1%), ciprofloxacin (97.1%) and cefixime (96.1%), followed by azithromycin 487 (71.6%) and ceftriaxone (33.3%). These results reflect the complexity of the resistance 488 mechanisms for azithromycin and ceftriaxone, where the known genetic determinants explain 489 only part of the antimicrobial susceptibility. However, specificity values (true negative rates, 490 TN/(TN+FP); TN=True Negatives, FP=False Positives) for these two antibiotics as well as 491 ciprofloxacin were above 99% (Additional file 1: Table S3), demonstrating that the genetic 492 mechanisms included in the database have a role in AMR. The specificity value for cefixime was 493 lower but nearly 90%, mainly due to the high number of isolates with an MIC below the threshold 494 but with three mutations characterising a mosaic penA allele (G545S, I312M and V316T, TP=367, 495 TN=323, PPV=53.2%; Additional file 1: Table S4). Benzylpenicillin and tetracycline showed 496 specificity values of 77.3% and 61.3%, respectively. In the first case, all the mechanisms included in the library showed a PPV value above 94%. For tetracycline, a considerable number of false 497 498 positive results are mainly caused by the presence of rpsJ V57M, for which PPV=83.8% 499 (TP=1083, FP=209; Additional file 1: Table S4). However, this mutation was kept in the AMR 500 library because it can cause intermediate resistance to tetracycline on its own (Additional file 3: 501 Figure S9).

Results from the benchmark analysis on the 3,987-isolates dataset were used to curate and optimize the AMR library. Thus, in order to objectively validate it, the benchmark analysis was also run on a combination of three different collections (N=1,607, Additional file 1: Table S1) with available MIC information for seven antibiotics including spectinomycin (Additional file 1: Table S3) (69, 70, 117). Results from the test and validation benchmark runs were compared, showing that sensitivity values on the six overlapping antibiotics were very similar, with the validation benchmark performing even better for azithromycin and ceftriaxone (Figure 3c). In terms of

509 specificity, both datasets performed equally well for all antibiotics except for benzylpenicillin, in 510 which specificity drops in the validation benchmark. This is due to the penA ins346D mutation (TP=1125, FP=83) and the *blaTEM* genes (TP=525, FP=36), which despite showing false 511 512 positives, have a PPV above 93% (Additional file 1: Table S5). In general, discrepancies found 513 between the test and the validation benchmarks can be explained by particular mechanisms that 514 on their own show high predictive values and affect antibiotics for which we do not currently understand all the factors involved in resistance, such as azithromycin and the ESCs (Additional 515 516 file 1: Table S5).

### 517 Over 12,000 public genomes available

518 Data for 11,461 isolates were successfully assembled and passed all quality cut-offs, resulting in 519 12,515 isolates after including the previously-available Euro-GASP 2013 dataset (15). New 520 assemblies were uploaded and made public on Pathogenwatch, which now constitutes the largest 521 repository of curated *N. gonorrhoeae* genomic data with associated metadata, typing and AMR 522 information at the time of submission of this manuscript. Updated data spans 27 different 523 publications (18, 53, 58-61, 63-65, 67-70, 117-131) and is organized into individual collections 524 associated with the different studies (Additional file 1: Table S6). Available metadata was added 525 for the genomes from these publications while basic metadata fields were kept for others (country, 526 year/date and ENA project number).

527 We cross-checked that the main clusters found in the phylogenetic trees obtained after creating 528 the public collections in Pathogenwatch were consistent with those observed in the trees in the 529 corresponding publications. For example, recent works defined two major clusters of N. 530 gonorrhoeae, termed Lineages A and B, which were found to be consistent with the corresponding 531 Pathogenwatch trees as exemplified for isolates from England in Town et al (2020) (68) (Figure 532 S11a). We were also able to differentiate the cefixime-resistant penA10 and penA34-carrying 533 clones from Vietnam from Lan et al (2020) (124) (Figure S11b) as well as the 10 major clusters 534 defined in the *N. gonorrhoeae* population circulating in New York City (NYC) as described in

535 Mortimer *et al* (2020) (120) (Figure S11c). In the last case, we also liked to emphasize the 536 usefulness of Microreact (37) as a parallel tool to Pathogenwatch for more complex visualization 537 purposes, such as showing the 10 major clusters in NYC as metadata blocks of different colours.

538 The *N. gonorrhoeae* public data available on Pathogenwatch spans nearly a century (1928-2018) 539 and almost 70 different countries (Additional file 3: Figure S12). However, sequencing efforts are 540 unevenly distributed around the world, and over 90% of the published isolates were isolated in only 10 countries, headed by the United Kingdom (N=3,476), the United States (N=2,774) and 541 542 Australia (N=2,388) (Additional file 1: Table S7, Figure 4). A total of 554 MLST, 1,670 NG-MAST 543 and 1,769 NG-STAR different STs were found in the whole dataset, from which a considerable 544 number were new profiles caused by previously undetected alleles or new combinations of known 545 alleles (N=92 new MLST STs, N=769 new NG-STAR STs and N=2,289 isolates with new NG-546 MAST porB and/or tbpB alleles). These new alleles and profiles were submitted to the 547 corresponding scheme servers.



Figure 4. Summary of the geolocalization and collection date of 12,515 public *N. gonorrhoeae* genomes in Pathogenwatch. Coloured bars represent the genotypic antimicrobial resistance (AMR) prediction based on the mechanisms included in the library. AZM = Azithromycin, CFM = Cefixime, CIP = Ciprofloxacin, CRO = Ceftriaxone, PEN = Benzylpenicillin, TET = Tetracycline.

548

549 Genomic studies are often biased towards AMR isolates, and this is reflected in the most 550 abundant STs found for the three typing schemes within the public data. Isolates with MLST 551 ST1901, ST9363 and ST7363, which contain resistance mechanisms to almost every antibiotic 552 included in the study, represent over 25% of the data (Figure 5). Isolates with MLST ST1901 and ST7363 are almost always associated with resistance to tetracycline, sulfonamides, 553 554 benzylpenicillin and ciprofloxacin and nearly 50% of isolates from these two types harbour 555 resistance mechanisms to cefixime. Ciprofloxacin resistance is not widespread among ST9363 556 isolates, which are associated with azithromycin resistance in nearly 50% of the isolates for this

557 ST (Figure 5). NG-STAR ST63 (carrying the non-mosaic penA-2 allele, penA A517G and mtrR 558 A39T mutations as described in (52)) is the most represented in the dataset and carries resistance 559 mechanisms to tetracycline, sulfonamides, and benzylpenicillin, but is largely susceptible to 560 spectinomycin, ciprofloxacin, the ESCs cefixime and ceftriaxone and azithromycin. NG-STAR 561 ST90 isolates, conversely, are largely associated with resistance to cefixime, ciprofloxacin and benzylpenicillin as they carry the key resistance mutations in mosaic penA-34, as well as in the 562 563 mtrR promoter, porB1b, ponA, gyrA and parC (as described in (52)). NG-MAST ST1407 is 564 commonly associated with MLST ST1901 and is the second most represented ST in the dataset 565 following NG-MAST ST2992, which mainly harbours resistance to tetracycline, benzylpenicillin 566 and sulfonamides (Figure 5).



Figure 5. Predicted antimicrobial resistance (AMR) profiles of the top five Multi-Locus Sequence Typing (MLST), *N. gonorrhoeae* Sequence Typing for Antimicrobial Resistance (NG-STAR) and *N. gonorrhoeae* Multi-Antigen Sequence Typing (NG-MAST) types in the *N. gonorrhoeae* public data in Pathogenwatch. Coloured circles in the grids show the proportion of genomes from each ST which are predicted to have an intermediate (susceptible but increased exposure) or resistant phenotype, red) versus susceptible genomes (in dark blue) from each sequence type (ST) and antibiotic. Bars on the top show the number of isolates from each ST coloured by the number of antibiotics the genomes are predicted to be resistant to.

#### 567 Case study: global expansion of an mtr mosaic-carrying clone

568 The genetic mechanisms that have commonly been associated with an increased MIC of 569 azithromycin in N. gonorrhoeae are two mutations in the 23S rRNA gene (A2045G and C2597T 570 substitutions, in *N. gonorrhoeae* nomenclature) as well as mutations in *mtrR* and its promoter 571 (132, 133). As described above, other mechanisms have also been recently discovered that 572 increase the MIC of azithromycin (Table 2), such as mosaicism affecting the efflux pump-encoding 573 *mtrCDE* genes and its repressor *mtrR*, mainly when the mosaic spans the *mtrR* promoter region 574 and mtrD gene (23, 24). Some studies have recently reported the local expansion of azithromycin-575 resistant N. gonorrhoeae lineages carrying an mtr mosaic in the USA (122, 123, 134) and 576 Australia (118). However, the extent of the dispersion of this mechanism to other parts of the 577 world has not been studied yet. Here, using the public genomes of N. gonorrhoeae in 578 Pathogenwatch, we have been able to explore this question.

579 A total of 1,142 strains with genetic determinants of azithromycin resistance were selected in 580 Pathogenwatch and combined with 395 genomes from a global collection (64) for background 581 contextualization (see Pathogenwatch project in (135)) (Figure 6a). 571 of the strains predicted 582 to be resistant to azithromycin had some form of mosaic in the *mtrR* promoter and/or *mtrD* gene 583 of one of the three types described in Wadsworth et al. (2018) (23) and included in the 584 Pathogenwatch AMR library (Table 2). These mosaics have been experimentally proven to 585 increase MIC of azithromycin above 1 mg/L, which is the EUCAST ECOFF value as well as the 586 Clinical Laboratory and Standards Institute (CLSI) non-susceptibility breakpoint (23, 24). One of 587 the *N. lactamica*-like mosaics, termed here '*mtr* mosaic.2', was by far the most extended, as it 588 was found in 545 genomes spanning the *mtrR* promoter and/or the *mtrD* gene, with 521 (95.6%) 589 of them spanning both regions. Twenty-five genomes contained a N. meningitidis-like mosaic 590 mtrR promoter and/or mtrD gene ('mtr mosaic.1') and in only 9 (36%) of them the mosaic 591 spanned both loci. The N. lactamica-like 'mtr mosaic.3' was only found in isolate ERR855360 592 (GCGS834) from Los Angeles (USA, 2012), which is where the reference sequence for this 593 mosaic was extracted from. Of the studies where these *mtr* mosaic-carrying genomes were 594 obtained from, only those from the USA and Australia specifically targeted and found this genetic 595 determinant of resistance. The rest did not target this mosaic and some of them found strains with 596 unexplained increased MICs of azithromycin (69, 121, 129), which could partly be explained by 597 the presence of these *mtr* mosaics.

598 We observed one main lineage carrying mosaic 2 in *mtrR* promoter and *mtrD* gene (Figure 6a) 599 with 520 genomes. Of those, only 3 and 8 isolates carried the 23S rDNA A2045G and C2597T 600 mutations, respectively. Interestingly, the first strain in the database with this type of mosaic dates 601 from 2006 (18), however, it was not until the end of 2011-2012 when this lineage started to expand 602 (Figure 6b). Despite the genomic data contained in Pathogenwatch being biased to the amount 603 of data sequenced and published from each country and year, we can easily infer that this lineage 604 has spread across the world as we detect cases in Australia (n=293) (118), the USA (n=195) (18, 605 120, 122, 123), Norway (n=19) (121), the United Kingdom (n=11) (68, 119), and Ireland (n=3) 606 (129). A strong association was found to the country of isolation (Figure 6c), with a broad diversity 607 of sublineages having spread across the USA (strains mostly isolated between 2012 and 2016). 608 In contrast, an expansion of a particular clone, likely from a single main introduction, was observed 609 to have occurred in Australia (strains isolated in 2017), followed by a further divergence of a 610 subclone within the country which correlates with the loss of the porB1b G120K and A121N 611 mutations (Figure 6d), likely through a recombination event. Despite epidemiological data not 612 being available for the Australian study (118), from their work we know that the clusters carrying 613 an mtr mosaic were mostly linked to transmission between men, although bridging among MSM 614 and heterosexual populations was also observed.

The results from our case study show that there is an emerging lineage of *N. gonorrhoeae* that has spread across the world and that is carrying a mosaic *mtr* that has been associated with lowto-medium resistance to azithromycin. This global lineage, as well as others that may emerge carrying this or other genetic AMR mechanisms, has to be closely monitored. For this purpose, an up-to-date genomic epidemiology tool such as Pathogenwatch, which includes a list of genetic

620 AMR mechanisms approved by an expert group is a great resource for the scientific community. 621 At the moment, Pathogenwatch includes references for three types of mosaics in the mtrR 622 promoter and *mtrD* genes that have been experimentally proven to increase MIC of azithromycin 623 (23, 24), and the detection of these mosaics on new genomes respond to a set of similarity rules 624 (see Data availability section). However, we will keep the database updated with new 625 experimentally-confirmed reference sequences that may arise from further studies as it is still 626 unclear whether all mosaics affecting the mtrCDE efflux pump will cause a decreased 627 susceptibility to azithromycin.



642 Figure 6. N. gonorrhoeae genomes carrying genetic AMR mechanisms associated to azithromycin resistance were 643 selected in Pathogenwatch (n=1,142) and combined with genomes from a global collection (64, 88) (total n=1528) for 644 background contextualization. (a) Main layout of the combined collection, with the emerging lineage carrying mtr mosaic 645 2 spanning the *mtrR* promoter and *mtrD* marked in red in the tree and the map. (b) Timeline of the genomes carrying 646 mtr mosaic 2 (in red) and other public genomes in the database without this genetic AMR mechanism. (c) Visualization 647 of the mtr mosaic 2-carrying lineage (n=520) spreading in the USA and Australia (see legend) using Microreact. The 648 arrow in turguoise colour marks the divergence of the Australian lineage, shown in more detail in (d) coloured by the 649 presence (in red) or absence (in white) of the porB1b G120K and A121N mutations. The Pathogenwatch project of this 650 case study can be explored in (135).

651

#### 652 Discussion

653 We present a public health focussed N. gonorrhoeae framework at Pathogenwatch, an open access platform for genomic surveillance supported by an expert group that can be adapted to 654 655 any public health or microbiology laboratory. Little bioinformatics expertise is required, and users 656 can choose to either upload raw short read data or assembled genomes. In both cases, the upload 657 of high-quality data is encouraged in the form of quality-checked reads and/or quality-checked 658 assemblies. Recent benchmark analyses show particular recommendations for long-read or 659 hybrid data (136) as well as short read-only data (40, 137). On upload, several analyses are run 660 on the genomes, and results for the three main typing schemes (MLST, NG-MAST and NG-STAR) 661 as well as the detection of genetic determinants of AMR and a prediction of phenotypic resistance 662 using these mechanisms can be obtained simultaneously. The library of AMR determinants 663 contained in Pathogenwatch for N. gonorrhoeae has been revised and extended to include the 664 latest mechanisms and epistatic interactions with experimental evidence of decreasing 665 susceptibility or increasing resistance to at least one of eight antibiotics (Tables 2). A test and 666 validation benchmark analyses revealed sensitivity and/or specificity values >90% for most of the 667 tested antibiotics (Additional file 1: Table S3). Sensitivity values for the antimicrobials in the 668 current dual treatment, azithromycin (80%) and ceftriaxone (50%), reflect the complexity of the 669 resistance mechanisms for these antibiotics, for which we can only explain part of the observed 670 phenotypic resistance. However, their specificity values were above 99% (Additional file 1: Table 671 S3), further strengthening the associations of the included AMR determinants in increasing MICs

of these antibiotics. It remains essential to perform phenotypic susceptibility testing so we can detect inconsistencies between phenotypic and genotypic data that can lead to the identification and subsequent verification of novel or unknown resistance mechanisms. This will allow to continuously expand the list of genetic AMR mechanisms, and the AMR prediction from genomic data will further improve.

677 The continuous increase in reporting of *N. gonorrhoeae* AMR isolates worldwide led to a call for 678 international collaborative action in 2017 to join efforts towards a global surveillance scheme. This 679 was part of the WHO global health sector strategy on STIs (2016-2021), which set the goal of 680 ending STI epidemics as a public health concern by year 2030 (7, 8). Several programmes are 681 currently in place at different global, regional or national levels to monitor gonococcal AMR trends, 682 emerging resistances and refine treatment guidelines and public health policies. This is the case 683 of, for example, the WHO Global Gonococcal Antimicrobial Surveillance Programme (WHO 684 GASP) (7, 8), the Euro-GASP in Europe (6, 15, 138), the Gonococcal Isolate Surveillance Project 685 (GISP) in the United States (139), the Canadian Gonococcal Antimicrobial Surveillance 686 Programme (140), the Gonococcal Surveillance Programme (AGSP) in Australia (141) or the 687 Gonococcal Resistance to Antimicrobials Surveillance Programme (GRASP) in England and 688 Wales (142). The WHO in collaboration with CDC has recently started an enhanced GASP 689 (EGASP) (143) in some sentinel countries such as the Philippines and Thailand (144), aimed at 690 collecting standardized and quality-assured epidemiological, clinical, microbiological and AMR 691 data. On top of these programs, WHO launched the Global AMR Surveillance System (GLASS) 692 in 2015 to foster national surveillance systems and enable standardized, comparable and 693 validated AMR data on priority human bacterial pathogens (145). Efforts are now underway to link 694 WHO GASP to GLASS. However, gonococcal AMR surveillance is still suboptimal or even lacking 695 in many locations, especially in LMICs, such as several parts of Asia, Central and Latin America, 696 Eastern Europe and Africa, which worryingly have the greatest incidence of gonorrhoea (3). 697 LMICs often have access to antimicrobials without prescription, have limited access to an optimal 698 treatment, lack the capacity needed to perform a laboratory diagnosis due to limited or nonexistent quality-assured laboratories, microbiological and bioinformatics expertise or training,
insufficient availability and exorbitant prices of some reagents on top of a lack of funding, which
altogether compromises infection control.

702 High throughput sequencing approaches have proved invaluable over traditional molecular 703 methods to identify AMR clones of bacterial pathogens, outbreaks, transmission networks and 704 national and international spread among others (28, 29). Genomic surveillance efforts to capture 705 the local and international spread of *N. gonorrhoeae* have resulted in several publications within 706 the last decade involving high throughput sequence data of thousands of isolates from many 707 locations across the world. The analysis of this data requires expertise, not always completely 708 available, in bioinformatics, genomics, genetics, AMR, phylogenetics, epidemiology, etc. For 709 lower-resourced settings, initiatives such as the NIHR Global Health Research Unit, Genomic 710 Surveillance of Antimicrobial Resistance (146) are essential to build genomic surveillance 711 capacity and provide the necessary microbiology and bioinformatics training for quality-assured 712 genomic surveillance of AMR.

713 One of the strengths of genomic epidemiology is being able to compare new genomes with 714 existing data from a broader geographical level, which provides additional information on, e.g. if 715 new cases are part of a single clonal expansion or multiple introductions from outside a specific 716 location. To support this, Pathogenwatch calculates phylogenetic trees from a set of genomes 717 selected as collections. Currently, over 12,000 isolates of N. gonorrhoeae have been sequenced 718 using high throughput approaches and publicly deposited on the ENA linked to a scientific 719 publication. We have quality-checked and assembled these data using a common pipeline and 720 we made it available through Pathogenwatch, with the aim of representing as much genomic 721 diversity of this pathogen as possible to serve as background for new analyses. These public 722 genomes are associated with at least 27 different scientific publications, and have been organized 723 in Pathogenwatch as individual collections (Additional file 1: Table S6). The clustering of strains 724 on the resulting reconstructions was found consistent with those in the original publications (some

examples in Figure S11), while differences in branch lengths may be attributed to the usage ofdifferent reconstruction methods.

727 The power of Pathogenwatch to investigate questions of public health concern is reflected in a 728 case study (Figure 6). By selecting 1,142 azithromycin resistant strains from the public data in 729 Pathogenwatch in the context of a global collection (64), we observed one clone carrying N. 730 lactamica-like mtr mosaic ('mosaic 2') in both the mtrR promoter and mtrD genes, likely resulting 731 from the same recombination event. Strong geographical structure was found in these 732 azithromycin resistant strains, with isolates from the USA (mostly from 2012-2016) clearly 733 differentiated from those from Australia (from 2017), which show a more clonal dispersion, likely 734 from a single main introduction to the country followed by a rapid spread. Interestingly, a 735 sublineage of this Australian mtr mosaic-carrying clone seems to have also diverged after losing the porB1b G120K and D121N mutations. It is important to note that the data from which these 736 737 inferences were derived was gathered from surveillance-based studies and outbreak 738 investigations, which may bias the observed global diversity of strains carrying this mosaic. 739 Phenotypic susceptibility data for azithromycin or epidemiological information were not available 740 for over half of these strains, thus impeding making further inferences. This reflects the need of 741 improving the submission of anonymized epidemiological and antimicrobial susceptibility data for 742 individual isolates rather than aggregated data to public repositories and/or as supplementary 743 information of the corresponding publications, as this is where the public data in Pathogenwatch 744 is coming from.

In this study, we have additionally gathered an advisory group of *N. gonorrhoeae* experts in different fields such as AMR, microbiology, genetics, genomics, epidemiology and public health who will consult and discuss current and future analytics to be included to address the global public health needs of the community. We suggest this strategy as a role model for other pathogens in this and other genomic surveillance platforms, so the end user, who may not have full computational experience in some cases, can be confident that the analytics and databases underlying this tool are appropriate, and can have access to all the results provided by 752 Pathogenwatch through uploading the data via a web browser. We are aware that this is a 753 constantly moving field and analytics will be expanded and updated in the future. These updates 754 will be discussed within an advisory group to make sure they are useful in the field and the way 755 results are reported is of use to different profiles (microbiologists, epidemiologists, public health 756 professionals, etc.). Future analytics that are under discussion include the automatic submission 757 of new MLST, NG-STAR and NG-MAST STs and alleles to the corresponding servers, e.g. 758 PubMLST (48) and the automatic submission of data to public archives such as the ENA. Inter-759 connectivity and comparability of results with PubMLST is of particular interest, as this database 760 has traditionally been the reference for *Neisseria* sequence typing and genomics and it is widely 761 used by the *N. gonorrhoeae* community. Plasmid and *tetM/blaTEM* subtyping as recently 762 described (147) will also be considered within the development roadmap of Pathogenwatch. 763 Including a separate library to automatically screen targets of potential interest for vaccine design 764 (148-150) as well as targets of new antibiotics currently in phase III clinical trials (i.e. zoliflodacin 765 (151) or gepotidacin (152)) can also be an interesting addition to the scheme. Regarding AMR, 766 new methods for phenotypic prediction using genetic data are continuously being reported (62, 767 153, 154), especially those based on machine learning algorithms (155), and will be considered 768 for future versions of the platform. The prediction of MIC values or ranges instead of SIR 769 categories will allow users to decide whether to use EUCAST (156) or CLSI (157) guidelines for 770 categorization.

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#### 772 Conclusions

In summary, we present a genomic surveillance platform adapted to *N. gonorrhoeae*, one of the main public health priorities compromising the control of AMR infections, where decisions on existing and updated databases and analytics as well as how results are reported will be discussed with an advisory board of experts in different public health areas. This will allow scientists from both higher or lower resourced settings with different capacities regarding high

throughput sequencing, bioinformatics and data interpretation, to be able to use a reproducible and quality-assured platform where analyse and contextualise genomic data resulting from the investigation of treatment failures, outbreaks, transmission chains and networks at different regional scales. This open access and reproducible platform constitutes one step further into an international collaborative effort where countries can keep ownership of their data in line with national STI and AMR surveillance and control programs while aligning with global strategies for a joint action towards battling AMR *N. gonorrhoeae*.

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### 786 List of abbreviations

- 787 AGSP: Australian Gonococcal Surveillance Programme
- 788 AMR: Antimicrobial Resistance
- 789 AZM: Azithromycin
- 790 CDC: Centers for Disease Control and Prevention
- 791 CFM: Cefixime
- 792 cgMLST: Core Genome Multi-Locus Sequence Typing
- 793 CIP: Ciprofloxacin
- 794 CLSI: Clinical Laboratory and Standards Institute
- 795 CRO: Ceftriaxone
- 796 ECOFF: Epidemiological Cut-Off
- 797 EGASP: Enhanced Gonococcal Antimicrobial Surveillance Programme
- 798 ENA: European Nucleotide Archive
- 799 ESCs: Extended Spectrum Cephalosporins
- 800 EUCAST: European Committee on Antimicrobial Susceptibility Testing
- 801 Euro-GASP: European Gonococcal Antimicrobial Surveillance Programme
- 802 FN: False Negative
- 803 FP: False Positive
- 804 GASP: Gonococcal Antimicrobial Surveillance Programme

- 805 GISP: Gonococcal Isolate Surveillance Project
- 806 GRASP: Gonococcal Resistance to Antimicrobials Surveillance Programme
- 807 HIV: Human Immunodeficiency Virus
- 808 LMICs: Low and Middle-Income Countries
- 809 MIC: Minimum Inhibitory Concentration
- 810 MLST: Multi-Locus Sequence Typing
- 811 NG-MAST: N. gonorrhoeae Multi-Antigen Sequence Typing
- 812 NG-STAR: *N. gonorrhoeae* Sequence Typing for Antimicrobial Resistance
- 813 NPV: Negative Predictive Value
- 814 PEN: Benzylpenicillin
- 815 PPV: Positive Predictive Value
- 816 SNPs: Single Nucleotide Polymorphisms
- 817 ST: Sequence Type
- 818 STI: Sexually-Transmitted Infection
- 819 TET: Tetracycline
- 820 TN: True Negative
- 821 TP: True Positive
- 822 UK: United Kingdom
- 823 WGS: Whole Genome Sequencing
- 824 WHO: World Health Organization
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831	Declarations
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- 833 Ethics approval and consent to participate
- Not applicable.
- 835
- 836 Consent for publication
- 837 Not applicable.
- 838

#### 839 Availability of data and materials

The assemblies included in the current version of the *N. gonorrhoeae* Pathogenwatch scheme and used for the AMR benchmark analyses were generated from raw sequencing data stored in the ENA. Project accession numbers are included in Additional File 1: Tables S1 and S6. The generated assemblies can be downloaded from Pathogenwatch. The AMR library can be accessed from: <u>https://gitlab.com/cgps/pathogenwatch/amr-libraries/-/blob/master/485.toml</u>. The code to reproduce the figures and analyses in this manuscript can be found in <u>https://gitlab.com/cgps/pathogenwatch/publications/-/tree/master/ngonorrhoeae</u>.

847

### 848 Competing interests

849 The authors declare that they have no competing interests.

850

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879

#### 880 Authors' contributions

DMA conceived the Pathogenwatch application. CY, RG, KA, BT, AU and DMA developed the Pathogenwatch application. LSB and DMA contributed to the conception and design of the work. CY and LSB generated, updated and benchmarked the *N. gonorrhoeae* AMR library. BT, CY, AU and LSB obtained, quality-checked and reassembled the raw data from the ENA. LSB revised the assembled data, obtained all metadata available from the corresponding scientific publications and created collections. LSB, CY and DMA analysed the data. LSB and DMA drafted the

- manuscript. LSB, DMA, CY, SA, KCM, TDM, DG, MJC, YHG, IM, BHR, WMS, GS, KT, TW, SRH
- and MU contributed to the acquisition, technical and scientific interpretation and discussion of the
- data. LSB, DMA, MJC, YHG, IM, BHR, WMS, GS, KT, TW and MU agreed to participate in the *N*.
- 890 gonorrhoeae Pathogenwatch Scientific Steering Group before the preparation of this manuscript,
- and participated in virtual discussions. All authors read and approved the final manuscript.
- 892

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