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Modeling the effect of vaccination on selection for antibiotic resistance in Streptococcus pneumoniae

Citation for published version:

Davies, NG, Flasche, S, Jit, M & Atkins, KE 2021, 'Modeling the effect of vaccination on selection for antibiotic resistance in Streptococcus pneumoniae', Science Translational Medicine, vol. 13, no. 606. https://doi.org/10.1126/scitransImed.aaz8690

Digital Object Identifier (DOI):

10.1126/scitranslmed.aaz8690

Link:

Link to publication record in Edinburgh Research Explorer

Document Version: Peer reviewed version

Published In: Science Translational Medicine

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1	Stabilising effects of competition and diversity determine vaccine impact on
2	antibiotic resistance evolution
3	
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- 16
- 17 **One sentence summary:** Frequency-dependent competition and extrinsically-
- 18 maintained diversity shape selection for antibiotic resistance following vaccination.

19 Bacterial vaccines can protect recipients from contracting potentially antibiotic-20 resistant infections. But by altering the selective balance between sensitive and 21 resistant strains, vaccines may also suppress—or spread—antibiotic resistance 22 among unvaccinated individuals. Predicting the outcome requires knowing what 23 drives selection for resistance in bacterial pathogens, and in particular, what 24 maintains the circulation of both antibiotic-sensitive and resistant strains of 25 bacteria. Using mathematical modelling, we show that the frequency of penicillin 26 resistance in Streptococcus pneumoniae (pneumococcus) across 27 European 27 countries can be explained by between-host diversity in antibiotic use, heritable 28 diversity in pneumococcal carriage duration, or frequency-dependent selection 29 brought about by within-host competition between resistant and sensitive 30 strains. We use our calibrated models to predict the impact of non-serotype-31 specific pneumococcal vaccination upon the prevalence of carriage, incidence of 32 disease, and frequency of resistance for *S. pneumoniae*. We find that the relative 33 strength and directionality of competition between resistant and sensitive 34 pneumococcal strains is the most important determinant of whether vaccination 35 promotes, inhibits, or has little effect upon the evolution of antibiotic resistance. 36 Finally, we show that country-specific differences in pathogen transmission 37 substantially alter the predicted impact of vaccination, highlighting that policies 38 for managing resistance with vaccines must be tailored to a specific pathogen and 39 setting.

40

41 In an age of widespread antibiotic resistance, there is growing interest in using vaccines 42 to prevent bacterial infections that would otherwise call for treatment with antibiotics 43 (1-4). This interest arises for two main reasons: first, vaccines are effective against both 44 antibiotic-resistant and antibiotic-sensitive bacteria; and second, successful prophylaxis 45 removes the need for a course of antibiotic therapy that might promote more resistance 46 (2–5). Over the past two decades, the use of pneumococcal conjugate vaccines (PCVs) 47 has seemingly borne out these advantages. Administering PCVs to young children has 48 substantially reduced disease caused by *S. pneumoniae* (5–8)—a common asymptomatic 49 coloniser of the nasopharynx which can cause pneumonia, meningitis and other 50 infections when invasive—and has decreased demand for antibiotic therapy, largely by 51 reducing cases of otitis media (5, 9). But because PCV formulations target only a fraction

52 of the ~100 known pneumococcal serotypes, the niche vacated by PCV-targeted 53 serotypes has been filled by non-vaccine serotypes, and overall pneumococcal carriage 54 has rebounded to pre-vaccine levels (10, 11). Concomitantly, the incidence of infections 55 attributed to non-vaccine serotypes (12) and the proportion of non-vaccine-type 56 infections exhibiting antibiotic resistance (5, 13) have risen in many settings. Concern 57 over serotype replacement—along with the high cost of PCV manufacturing—has spurred the development of "universal" (non-serotype-specific) whole-cell or protein-58 59 based pneumococcal vaccines protecting against all serotypes, some of which are now 60 in early-stage clinical trials (14). If successful, universal pneumococcal vaccines could 61 reduce the burden of pneumococcal disease without selecting for serotype replacement. 62

63 However, it is unclear how universal vaccination itself may impact upon the evolution of 64 antibiotic resistance in *S. pneumoniae*, which is a concern given that vaccination is 65 unlikely to eliminate pneumococcal carriage entirely (15). Mathematical models of 66 bacterial transmission can be used to predict the impact of vaccination on antibiotic 67 resistance (16, 17), but existing models for *S. pneumoniae* focus on serotype-specific 68 vaccines and, even then, disagree over the expected impact of vaccination on resistance 69 evolution (18-24). Comparing and interpreting the results of these models is hampered 70 by the fact that none starts from a position of recapitulating large-scale empirical 71 patterns of antibiotic resistance. The main challenge in replicating these patterns lies in 72 identifying the mechanisms that maintain long-term coexistence between sensitive and 73 resistant pneumococcal strains across a wide range of antibiotic treatment rates, like 74 those seen across Europe and the United States (25, 26). Robust predictions of the long-75 term impact of non-serotype-specific vaccination on resistant pneumococcal disease 76 require a mechanistic understanding of these patterns. 77

78

Results

79

80 Stability in resistance evolution can be maintained by frequency-dependent

81 **competition or extrinsically-imposed diversity.** A model must be able to explain the

- 82 current burden of an infectious disease before it can be used to robustly predict the
- 83 impact of interventions for managing that disease. Across Europe, the frequency of
- 84 antibiotic resistance among isolates from pneumococcal infections shows two salient

85 features for models to recapitulate (Fig. S1). One feature is spatial: the frequency of 86 penicillin non-susceptibility varies between countries, and is higher in countries where 87 more penicillin is consumed (27). The other is temporal: although in individual 88 countries, resistance fluctuates from year to year, the overall frequency across Europe 89 of penicillin non-susceptibility in pneumococcal isolates has remained steady at roughly 90 12% since consolidated records began in 2005 (28). These observations contradict 91 simple models of resistance evolution, which predict that intermediate frequencies of 92 resistance cannot be stably maintained in the long term: that is, either sensitive strains 93 will competitively exclude resistant strains, or resistant strains will competitively 94 exclude sensitive strains, unless there is some mechanism that maintains coexistence 95 between them (25, 29).

96

97 By conducting a literature search, we identified nine such mechanisms (25, 26, 30–41) 98 that fall into two broad classes. In one class, coexistence is maintained by 99 environmental or genetic diversity that effectively creates separate niches for resistant 100 and sensitive strains, preventing them from completely overlapping in competition. In 101 the other class, competition between resistant and sensitive strains is itself the 102 stabilising factor that maintains coexistence, because resistant and sensitive strains 103 exhibit alternative competitive phenotypes that afford strains a competitive advantage 104 when rare, thus promoting negative frequency-dependent selection for resistance. Thus, 105 extrinsically-imposed diversity and frequency-dependent competition are two key 106 forces maintaining stability in resistance evolution. We find that four of the nine 107 identified mechanisms for maintaining coexistence are biologically plausible for S. 108 pneumoniae (Table 1).

109

110 Four models of resistance evolution. To compare these four mechanisms, we embed 111 each in a shared model framework of person-to-person transmission of nasopharyngeal pneumococcal carriage. This framework tracks the country-specific frequency of 112 113 resistance in pneumococci circulating among children under five years old, the age 114 group that drives the majority of pneumococcal transmission and disease (42, 43). We 115 assume that each individual makes effective contact with another random individual at 116 rate β , thereby potentially acquiring a strain (either sensitive or resistant) carried by 117 the contacted person. With probability c, resistant strains fail to transmit, where c

118 represents the transmission cost of resistance (44, 45). A carrier naturally clears all 119 strains at rate u, and is exposed to antibiotic therapy at a country-specific rate τ , which 120 clears the host of sensitive strains only. We assume this treatment rate is independent

- 121 of carriage status (46) and we do not explicitly track disease progression in hosts.
- 122

123 Under the "Treatment diversity" and "Pathogen diversity" models, extrinsically-124 maintained diversity among hosts or among pathogens prevents competitive exclusion 125 by keeping resistant and sensitive strains from fully competing with each other. In the 126 "Treatment diversity" model (Fig. 1a), heterogeneity in the consumption of antibiotics 127 between host subpopulations within a country maintains coexistence (25, 34, 35). These 128 subpopulations could correspond to geographical regions, socioeconomic strata, host 129 age and risk classes, or a combination of these. Provided that transmission between 130 high-consumption (resistance-promoting) and low-consumption (resistance-inhibiting) 131 subpopulations is not too frequent, an intermediate frequency of resistance can be 132 maintained across the whole population. Because coexistence is maintained by 133 assortative mixing between subpopulations differing in antibiotic use, the key 134 parameters governing coexistence in this model are κ, the variability in antibiotic 135 consumption between subpopulations, and *g*, the relative rate at which within-country 136 contact is made within subpopulations rather than between them (Fig. S2).

137

In the "Pathogen diversity" model (Fig. 1b), pneumococci are divided into subtypes ("D-138 139 types"(38)) that vary in their mean duration of natural carriage. All else equal, the D-140 type with the longest carriage duration would be expected to competitively exclude all 141 other strains; the model assumes that diversifying selection acting on the D-type locus 142 keeps all subtypes in circulation. What D-types correspond to is not explicitly specified 143 by this model, but one candidate is serotype variation. For example, if antigenic 144 diversity is promoted by host acquired immunity to capsular serotypes, and serotypes 145 tend to differ in their intrinsic ability to evade clearance by the immune system, then 146 intermediate resistance can be maintained because selection for resistance tends to be 147 greater in serotypes that have a longer duration of carriage (38). Long-lasting serotypes 148 will tend to evolve resistance, while shorter-lived serotypes will tend not to—a pattern 149 observed in *S. pneumoniae* (38) and reproduced by this model (Fig. S3). The parameters

- 150 governing coexistence in this model are *a*, the strength of diversifying selection on the
- 151 D-type locus, and δ , the variability between subtypes in clearance rate.
- 152

153 Under the "Treatment competition" and "Growth competition" models, coexistence is 154 maintained by within-host competition between sensitive and resistant strains. In these 155 models, hosts can be co-colonised by multiple strains. Then, competition between strains within the host niche determines which strain is transmitted to other potential 156 157 hosts (26). The "Treatment competition" model (Fig. 1c) assumes that antibiotic therapy 158 mediates within-host competition, such that when a co-colonised host takes antibiotics 159 (*i.e.*, at rate τ), the sensitive strains are cleared and only the resistant strains are 160 transmitted to other hosts. The "Growth competition" model (Fig. 1d) has both 161 treatment-mediated and growth-mediated competition: while in the presence of 162 antibiotics, resistant strains still outcompete co-colonising sensitive strains, in the 163 absence of antibiotics, sensitive strains gradually outcompete co-colonising resistant 164 strains at rate *b*. We assume that there is no transmission cost of resistance in this latter 165 model (*i.e.*, c = 0); instead, the within-host growth advantage *b* of sensitive strains 166 accounts for the cost of resistance. In these competition models, resistant strains have 167 an advantage in antibiotic-mediated competition, while sensitive strains have an advantage in growth-mediated competition. These alternative forms of within-host 168 169 competition can both promote coexistence because rare strains can more consistently 170 exploit a competitive advantage over common strains, thus creating negative frequency-171 dependent selection for resistance (26). The key parameter governing coexistence in 172 these two models is *k*, the relative rate of co-colonisation compared to primary 173 colonisation.

174

175 In all four models, we assume that contact between individuals is assortative by 176 country, such that with probability *f*, contact is with a random person from the same 177 country, and with probability 1 – *f*, contact is with a random person from any country. 178 We implement these models using systems of ordinary differential equations. All four 179 models (25, 26, 38) are structurally neutral (25, 29), meaning that any coexistence 180 exhibited by the models is accounted for by the specified biological mechanism rather than by any bias in the logical structure of the model that generates coexistence "for 181 182 free" (29). Additionally, while the within-host competition models capture co-

- 183 colonisation using a simplified subset of only 2 "mixed-carriage" states (S_R and R_S, Fig.
- 184 1a&b), we have previously shown (26) that this is equivalent to a more complex
- 185 individual-based model with an arbitrary number of mixed-carriage states.
- 186

187 All four models reproduce observed patterns of resistance. The European Centre 188 for Disease Prevention and Control (ECDC) monitors antibiotic consumption and resistance evolution across European countries (13, 28). These data capture a natural 189 190 experiment in resistance evolution: for each monitored drug and pathogen, each 191 country reports a different rate of antibiotic consumption in the community and 192 exhibits a different frequency of resistance among invasive bacterial isolates. By fitting 193 models to this multi-country data set, we can potentially rule out models that cannot 194 reproduce the large-scale patterns that are observed. We use Bayesian inference to fit 195 the model-predicted equilibrium frequency of resistance to the reported frequency of 196 penicillin non-susceptibility in *S. pneumoniae* across 27 European countries, assuming a 197 50% carriage prevalence (11, 42) and a carriage duration of 47 days (47, 48) in children 198 under five years old. We begin by assuming that countries only differ by their reported 199 treatment rate—where we define a treatment course as equivalent to z = 5 defined daily 200 doses of penicillin—with other model parameters shared across countries.

201

Strikingly, each model fits equally well to the empirical relationship between resistance
and antibiotic use (all model WAICs are similar; Fig. 2a) and recovers plausible
posterior parameter distributions (Fig. 2b; Fig. S4). That is, the empirical data do not
distinguish between the four alternative mechanisms of resistance evolution we have
identified. Later, we relax the assumption that only the treatment rate varies between
countries, allowing us to capture additional between-country variation in resistance not
explained by population-wide penicillin consumption.

209

210 Mechanisms of resistance evolution determine the impact of vaccination on

211 **resistant disease.** To determine the impact of universal vaccination on pneumococcal

212 disease, we consider three outcomes. The first is the impact of the vaccine upon the

- 213 prevalence of pneumococcal carriage. The second is the vaccine impact upon the
- 214 frequency of penicillin resistance among circulating pneumococcal strains remaining
- 215 after vaccination. The third is the impact of the vaccine upon the prevalence of resistant

216 pneumococcal carriage—*i.e.*, the prevalence of carriage multiplied by the frequency of 217 penicillin resistance. Since all four models are equally capable of recapitulating 218 observed patterns of penicillin resistance in *S. pneumoniae*, our aim is to determine 219 whether the mechanism maintaining stability in resistance evolution—frequency-220 dependent competition or extrinsically-imposed diversity—matters when forecasting

- the impact of interventions for managing resistance.
- 222

223 We consider two alternative non-serotype-specific vaccines: an "acquisition-blocking" 224 vaccine, which prevents carriage from being established with probability ε_a , and a 225 "clearance-accelerating" vaccine, which shortens the duration of carriage by a fraction 226 ε_{c} . Both vaccines reduce pneumococcal transmission through alternative modes of host 227 immunity that might be elicited by a whole-cell or protein-based universal 228 pneumococcal vaccine. Analogously to naturally-acquired serotype-independent 229 pneumococcal immunity (49), the protective effect of whole-cell vaccines manifests as 230 accelerated clearance (50); it is unclear whether protein-based vaccines would block 231 pneumococcal acquisition, like PCVs, or accelerate clearance (51). We refer to ε_a or ε_c as 232 the vaccine efficacy, and for simplicity, we assume that all children under five years old 233 have vaccine protection, as would be established by an infant vaccination programme 234 rolled out across Europe. In order to compare these vaccines with an alternative 235 intervention of antibiotic stewardship, we also evaluate the impact of reducing the rate 236 of penicillin prescribing by a fraction ε_s .

237

We find that both vaccines have a similar impact upon carriage prevalence, regardless of whether competition or diversity maintains stability in resistance evolution (Fig. 3a). Specifically, as the vaccine efficacy ε_a or ε_c increases, carriage decreases, with the elimination of pneumococcal carriage occurring at a vaccine efficacy between 50 and 60%. Reducing antibiotic prescribing moderately increases pneumococcal carriage, such that carriage prevalence increases to approximately 54% across all countries when penicillin prescribing is eliminated completely.

However, the mechanism of resistance evolution has a substantial impact upon whether
vaccines increase or decrease the frequency of resistance in *S. pneumoniae* in the long
term (Fig. 3b). In the "Treatment diversity" and "Pathogen diversity" models, the

249 acquisition-blocking vaccine has relatively little impact upon the frequency of 250 resistance, because administering a universal pneumococcal vaccine to all individuals 251 does not substantially alter the distribution of antibiotic use or of heritable variation in 252 clearance rates. By contrast, in the within-host competition models, vaccination has a 253 substantial impact upon resistance evolution because by reducing pneumococcal 254 circulation, vaccines decrease the rate at which strains encounter each other within 255 hosts, and hence strongly decrease competition between pneumococcal strains. 256 Specifically, the acquisition-blocking vaccine selects strongly against resistance in the 257 "Treatment competition" model: since antibiotic-mediated within-host competition 258 benefits the resistant strain in this model, the vaccine works against this competitive 259 advantage and therefore inhibits resistance. Conversely, in the "Growth competition" 260 model, growth-mediated competition benefits the sensitive strain, and so by reducing 261 competition, vaccination tends to promote resistance. These results expand upon our 262 previous finding that the rate of co-colonisation modulates resistance evolution through 263 its impact upon within-host competition (26).

264

265 The clearance-accelerating vaccine exhibits similarly divergent impacts across 266 mechanisms of resistance evolution. However, compared with the acquisition-blocking 267 vaccine, it also has an additional resistance-inhibiting effect across all models, because a 268 shorter duration of carriage—whether natural or vaccine-induced—selects against 269 resistance (38). This suggests that vaccines that accelerate natural clearance have a 270 particular potential for managing resistant infections. As expected, reducing the rate of 271 penicillin prescribing selects against resistance, exhibiting a similar impact across all 272 four models.

273

The impact on resistant carriage (Fig. 3c), which combines changes in the prevalence of
carriage and changes in the frequency of resistance, can be treated as a proxy for the
incidence of resistant infections. Overall, under the "Growth competition" model,
vaccination at intermediate efficacy is expected to increase the rate of resistant carriage,
and hence the number of cases of resistant disease. In other models, vaccination always
reduces resistant carriage, particularly under the "Treatment competition" model. A
summary of the strongest vaccine impacts is shown in Fig. 3d.

282 Evidence to inform policy and vaccine trials. For vaccines to be considered an 283 efficient means of controlling resistant infections, they must compare favourably to 284 existing interventions, such as reducing inappropriate antibiotic use (52). The UK 285 government has recently announced an initiative to reduce antibiotic consumption by 15% by the year 2024 (52). Our models predict that a 15% reduction in primary-care 286 287 penicillin consumption would reduce carriage of penicillin-non-susceptible 288 pneumococci from 6% to 3%. The vaccine efficacy required to yield the same effect 289 varies considerably depending upon the mechanism of resistance evolution (Fig 4a); for 290 example, the required vaccine efficacy is lowest under the "Treatment competition" 291 model ($\varepsilon_a = 11\%$ or $\varepsilon_c = 7\%$), and highest under the "Growth competition" model ($\varepsilon_a =$ 292 52% or ε_c = 50%). A full comparison of vaccine and stewardship interventions would 293 require accounting for the economic cost of vaccines versus antibiotics, the wider range 294 of resistant pathogens that would be targeted by restrictions on antibiotic use, and any 295 potential increase in pathogen circulation that might be brought about by inadvertent 296 decreases in appropriate antibiotic use.

297

298 In randomized controlled trials of pneumococcal vaccines, resistance-related endpoints 299 have routinely been evaluated over a follow-up period of between 6 months and 3.5 300 years after vaccination (53, 54). If vaccine-induced changes in resistance evolution 301 unfold over a considerably longer timescale, similarly-designed trials may not fully 302 capture vaccine impact on resistance. Indeed, we find that it can take 5–10 years for 303 resistance to stabilise following vaccination (Fig. 4b), and that short-term drops in 304 resistance can be reversed—or even give way to increased resistance—in the long term. 305 Moreover, a trial in which vaccination is not offered to a substantial fraction of the 306 population would not capture the full impact of reduced pneumococcal circulation, 307 which is what drives competition-mediated changes in resistance in our models. Finally, 308 our analysis assumes that vaccines are administered to all recipients simultaneously. In 309 the real world, where vaccination is likely to be be rolled out gradually, the full effect of 310 vaccination would take even longer to observe.

311

312 The impact of vaccination at a national level varies depending upon the treatment rate

313 in a given country. Focusing on the specific outcome of childhood pneumococcal

314 pneumonia cases, we find that while interventions have a consistent impact from

country to country on the total pneumonia case rate, the impact on resistant pneumonia
cases is greatest in those countries where antibiotic use, and hence resistance, is highest

- 317 (Fig. 4c). We focus on resistant carriage, but the realised public health benefits of any
- 318 intervention targeting both resistant and sensitive strains will depend upon the relative
- 319 health burdens of susceptible versus non-susceptible *S. pneumoniae* infections;
- 320 enumerating these comparative burdens is the subject of ongoing research (55).
- 321

322 Vaccination in a high-burden setting. High prevalences of carriage, disease, and 323 resistance are often observed in low-income settings, and it is desirable to know 324 whether this could substantially alter predictions of vaccine impact. As an illustrative 325 example, a 90% pneumococcal carriage rate, with 81% of isolates resistant to penicillin, 326 has been observed among children under five years old in western Kenya (56). This may 327 be partly attributable to a longer average duration of carriage in this setting, as a 71-day 328 mean duration of natural pneumococcal carriage has been measured in Kilifi, eastern 329 Kenya (57).

330

331 To model a similar high-burden setting, we adjust model parameters estimated from 332 European data: increasing the mean natural carriage duration, transmission rate, and 333 treatment rate to match observed data, and ignoring mixing with any other countries (*f* 334 = 1), while keeping other parameters the same. We find that a comparatively greater 335 vaccine efficacy is needed to reduce the prevalence of resistant carriage in a high-336 burden, high-resistance setting (Fig. 5). This is particularly true under the "Growth 337 competition" model, because in this model resistant carriage only declines as total 338 pneumococcal carriage declines, and it is particularly difficult to reduce overall carriage 339 in a high-transmission setting. Simultaneously, vaccination may have a comparatively 340 greater impact in high-burden settings because of a comparatively higher incidence of 341 disease: for example, Kenya has been estimated to have an 8.8-fold higher incidence of 342 severe pneumococcal pneumonia than the average in Europe (58).

343

344 Accounting for additional between-country variation does not substantially alter

345 **predictions.** Our focus thus far has been on the impact of the four identified

346 mechanisms *per se* upon resistance evolution, and accordingly we have focused on

347 reproducing the positive association between treatment rate and resistance frequency

348 rather than attempting to capture the additional variability in resistance frequency 349 between countries not accounted for by the reported treatment rate alone (Fig. 2a). This 350 additional variability may partially stem from differences in national testing and 351 reporting practices, or between-country differences in the distribution of pneumococcal 352 serotypes among invasive isolates (59). However, another possibility is that this 353 additional variability in resistance results from systematic differences in pathogen 354 biology or host behaviour across countries which can be captured by our modelling 355 framework.

356

357 To help identify which model parameters could account for this variability, we relax the 358 assumption that only the treatment rate varies across countries, and perform Bayesian 359 maximum *a posteriori* fitting, assuming one additional parameter (*c*, *b*, β , *u*, *f*, *z*, *g*, κ , *a*, δ , 360 or k) is free to vary between countries while other parameters are held constant. We 361 find that additional variation in resistance between countries can be explained by 362 variation in certain other parameters, depending upon which model is used (Fig. 6a-b). 363 Importantly, among those parameters for which additional variation between countries 364 can explain the variation in resistance (Fig. 6c), predictions for the overall impact of 365 vaccination remain similar, with the major differences between scenarios still 366 attributable to the underlying mechanism of resistance evolution (Fig. 6d; Figs. S5–S16). 367 Models that could make more accurate country-specific predictions would need to 368 account for the effects of demographic structure, differences in carriage prevalence and 369 disease rates between settings, and variable vaccine protection among individuals. 370 371 Discussion 372 373 We have identified four mechanisms of resistance evolution that are capable of

374 recapitulating the observed relationship between penicillin consumption and penicillin
375 non-susceptibility in *S. pneumoniae* across Europe. These mechanisms are not mutually
376 exclusive, but the relative importance of each is predicted to have a substantial impact
377 upon predictions for resistance evolution under vaccination. In particular, the
378 "directionality" of within-host competition—that is, whether, on average, within-host
379 competition tends to benefit resistant or sensitive strains—strongly determines
380 whether vaccination selects for a decrease or an increase in antibiotic resistance in the

- 381 long term. This directionality may vary between pathogens, but is also sensitive to the
- antibiotic treatment rate, and so may also vary between settings. Although we have
- 383 focused on competition between sensitive and resistant strains of *S. pneumoniae* only,
- 384 competition between serotypes (24) and with other bacteria colonizing the
- 385 nasopharynx will also impact upon resistance evolution, and determining the
- importance of these other sources of within-host competition is crucial.
- 387

388 A key result of our models is that the mode of vaccine protection—whether acquisition-389 blocking or clearance-accelerating—has an appreciable impact upon resistance 390 evolution. Whole-cell and purified-protein pneumococcal vaccines may induce 391 antibody-mediated humoral immunity, CD4+ T helper-17 cell-mediated immunity, or 392 both, with the type of immunity mediating pneumococcal acquisition, carriage, and 393 disease in ways that are still not fully understood (49–51). By modelling both modes of 394 vaccine action, we have highlighted that clearance-accelerating vaccines have increased 395 potential for preventing the spread of resistance, because in shortening the duration of 396 asymptomatic carriage they limit the fitness advantage of resistant pathogens under 397 selection pressure from antibiotic use.

398

399 We fit our models to a "snapshot" of penicillin non-susceptible S. pneumoniae as 400 observed across European countries in 2007, finding that each model recapitulated the 401 data equally well. This raises the question of what kind of data would be needed to 402 distinguish the models. One possibility would be to consider trends of resistance 403 evolution over time. Indeed, the prevalence of penicillin non-susceptibility in S. 404 pneumoniae remained largely stable in Europe between 2005–2017, a period which saw 405 the incorporation of PCV into the routine immunization schedules of most European 406 countries (60). This could be viewed as favouring the "diversity" mechanisms, which 407 predict little change in resistance evolution following vaccination. But because serotype 408 replacement has largely negated any vaccine impact on the prevalence of 409 nasopharyngeal pneumococcal carriage (10, 11), it is not clear that we would be able to 410 detect any effects of competition-mediated resistance evolution following a serotype-411 specific vaccine such as PCV—particularly given the complexity of detecting vaccine-412 attributable changes in resistance in a population-level associational study that would 413 be confounded both by serotype replacement and by other changes in resistance

evolution that might be expected to occur at a national level over the course of multiple
years. However, it is known that the prevalence of pneumococcal carriage declines
substantially with age (42). Therefore, it might be possible to detect a signal of withinhost competition between sensitive and resistant strains by comparing the relative
prevalence of resistant pneumococcal carriage in younger versus older hosts, provided
that other differences could be controlled for.

420

421 Under the "Treatment diversity" and "Pathogen diversity" models, we have argued that 422 universal pneumococcal vaccination will have little impact upon the long-term 423 evolution of antibiotic resistance because it does not change the sources of diversity 424 that modulate resistance evolution. Nonetheless, it is possible to target vaccines such 425 that this diversity is harnessed to manage resistance: high-resistance serotypes could 426 be targeted with a serotype-specific vaccine, or high-treatment subpopulations could be 427 targeted for vaccination in order to more effectively manage resistance. Indeed, 428 vaccination does have an additional inhibiting effect upon resistance in our models 429 because of the latter effect. This inhibition occurs because the vaccine has a relatively 430 greater impact upon transmission in populations where the prevalence of carriage is 431 already low, which in our models occur in countries or subpopulations with more 432 antibiotic consumption. Since these populations drive resistance more strongly, the 433 vaccine's comparatively greater impact in these populations tends to moderately inhibit 434 resistance overall. We note that while previous work (38) has suggested that resistance 435 evolution under a "Pathogen diversity" model results in a "stepped" resistance pattern 436 in which D-types are either fully sensitive or fully resistant at equilibrium, we find that 437 small amounts of mixing between populations can smooth out this pattern and allow 438 intermediate rates of resistance within subtypes (Fig. S2). Finally, while we have framed 439 "Treatment competition" and "Growth competition" as two distinct alternatives, they 440 can instead be viewed as endpoints on a continuum, with possible models of resistance evolution for which both c > 0 and b > 0 lying between them. The impact of vaccination 441 442 on resistance in such a model would depend upon the relative importance of treatment-443 mediated and growth-mediated competition.

444

This analysis has necessarily made simplifying assumptions. We have focused on
prevalence (the fraction of individuals who are carriers) rather than incidence (the rate

447 of new carriage episodes) of nasopharyngeal carriage in presenting our findings. There 448 is evidence that pneumococcal disease progression is more likely to occur shortly after 449 nasopharyngeal acquisition (61), suggesting that incidence may be more relevant than 450 prevalence for predicting disease outcomes. Of particular note, recent modelling work 451 has suggested that clearance-accelerating vaccines can increase rates of pneumococcal 452 acquisition, if extended carriage is protective against new acquisition (62). However, it is not obvious how to compare rates of carriage acquisition across the models examined 453 454 in this paper, particularly because co-colonisation is explicitly tracked in some but not 455 all models. More work is required to clarify the links between acquisition, carriage, and 456 disease across competing models of pneumococcal transmission. Additionally, we have 457 assumed that antibiotic treatment rates among pneumococcal carriers remains constant 458 after the introduction of a vaccine, even though treatment rates dropped in many 459 settings following PCV introduction (5, 9). However, for a universal pneumococcal 460 vaccine that reduces antibiotic treatment rates because it reduces carriage and thereby 461 prevents antibiotic-treatable disease, any reduction in treatment will only occur among 462 individuals who, because of vaccine protection, are not pneumococcal carriers, all else 463 being equal. It might then be expected that treatment rates in carriers would remain 464 equally high among those individuals for whom vaccine protection has failed—although 465 physicians may be less inclined to prescribe antibiotics for respiratory tract infection 466 more generally after the introduction of a new pneumococcal vaccine. Finally, we have 467 focused on modelling children under 5 years old only. We would not expect 468 incorporating age structure to lead to qualitatively different results, but age-related 469 maturation of the immune system has been shown to be important for maintaining the 470 circulation of pneumococcal serotypes (49) which we have abstracted here.

471

472 Our work helps resolve the question: What explains the persistent coexistence between 473 resistant and sensitive strains of *S. pneumoniae*? (25) by demonstrating that multiple 474 mechanisms are capable of explaining trends of resistance across European countries. 475 Since there is empirical support for within-host competition between sensitive and 476 resistant pathogen strains (63–66), heritable differences in the propensity for resistance 477 within species (38, 67), and within-country heterogeneity in antibiotic consumption 478 rates (68–70), all of these mechanisms likely contribute to this pattern. Our results 479 contextualize previous mathematical studies which have variously suggested that

480 serotype-specific vaccination may increase (24), decrease (22) or have no impact upon 481 (18) the frequency of resistance in *S. pneumoniae*. While the potential for vaccination to 482 promote resistance because of competition between sensitive and resistant strains has 483 been described previously (24), we have shown that vaccination can either promote or 484 inhibit resistance depending upon the directionality of within-host competition. While 485 vaccines targeting highly-resistant serotypes can decrease resistance (22), we have 486 shown that a serotype-independent vaccine promoting accelerated natural clearance 487 can decrease resistance across all circulating subtypes. And where single-population 488 models have found no long-term impact of vaccination on resistance frequency (18), we 489 have shown that in multi-population models, vaccination can inhibit resistance if it has 490 a larger impact in subpopulations that consume more antibiotics. The direction and 491 magnitude of this effect would depend upon variation in vaccine uptake, vaccine 492 efficacy, and pathogen transmission among subpopulations, and we have not 493 systematically explored this variation here. 494 495 A highly efficacious serotype-independent pneumococcal vaccine can indeed reduce the

overall burden of antibiotic-resistant pneumococcal infections. However, the long-term
effect upon resistance of a vaccine with intermediate efficacy is less certain, as vaccine
impact depends crucially upon the mechanisms that drive resistance evolution. Thus,
empirical investigation of pathogen competitive dynamics—and the impact of settingspecific factors on these dynamics—is needed to make accurate predictions of vaccine
impact on resistant infections.

- 502
- 503
- 504

Methods

505 *Study design.* This study comprises four parts: a literature search used to identify 506 plausible mechanisms through which coexistence can be maintained between sensitive 507 and resistant pneumococcal strains across a range of antibiotic treatment rates; a 508 mathematical modelling study embedding these mechanisms of resistance evolution in 509 four models of pneumococcal transmission; a Bayesian statistical analysis to fit these 510 models to empirically observed frequencies of penicillin non-susceptibility and 511 community penicillin consumption across 27 European countries for the year 2007; and 512 a vaccine impact analysis using these fitted models to forecast the impact of a universal

- 513 pneumococcal vaccine. We use data from 2007 because changes in pneumococcal
- 514 resistance reporting standards for some countries after this year hamper the between-
- 515 country comparability of data (71). Our objectives were to identify the mechanisms
- 516 potentially responsible for maintaining coexistence between resistant and sensitive
- 517 pneumococci in Europe, and to determine whether the impact of vaccination on the
- 518 evolution of resistance depends upon which mechanism is assumed to operate.
- 519
- 520 *Mechanisms driving resistance.* We searched PubMed using the terms: (AMR OR ABR OR 521 ((antimicrobial OR antibiotic) AND resist*)) AND ((model OR modelling OR modeling) 522 AND (dynamic* OR transmi* OR mathematical)) AND (coexist* OR intermediate). This 523 yielded 93 papers (Table S1). We included all papers containing a dynamic host-to-host 524 pathogen transmission model analysing both sensitive and resistant strains with stable 525 coexistence as an outcome of the model. From the 11 studies meeting these criteria, we 526 identified nine unique mechanisms, two of which correspond to alternative 527 parameterisations of a within-host competition model. We ruled out four mechanisms 528 because of implausibility or because previous work shows that the mechanism does not 529 bring about substantial coexistence, leaving four mechanisms (Table 1). 530

531 *Model framework.* We analyse the evolution of antibiotic resistance by tracking the 532 transmission of resistant and sensitive bacterial strains among hosts in a set of *M* 533 countries indexed by $m \in \{1, 2, ..., M\}$ using systems of ordinary differential equations. 534

- In a simple model, hosts can either be non-carriers (X), carriers of the sensitive strain
 (S), or carriers of the resistant strain (R). Omitting country-specific subscripts *m* for
 concision, model dynamics within a country are captured by
- 538

539
$$dS/dt = \lambda_{S}X - (u + \tau)S$$

540 $dR/dt = (1 - c)\lambda_{R}X - uR$
541 $X = 1 - S - R$, (1)

542

543 where λ_s is the force of infection of the sensitive strain, λ_R is the force of infection of the 544 resistant strain, *c* is the transmission cost of resistance, *u* is the rate of natural 545 clearance, and τ is the treatment rate. In this model, in a given country, the total 546 carriage of the sensitive strain is *S* and the total carriage of the resistant strain is *R*.
547 Force of infection terms are defined below; a summary of all model parameters can be
548 found in Table S2.

549

The "Treatment diversity" model extends the simple model (eq. 1) by structuring each country into multiple subpopulations that exhibit different rates of antibiotic treatment and make contact with each other at unequal rates (25, 34, 35, 72). In each country, we model *N* equally-sized representative subpopulations indexed by $i \in \{1, 2, ..., N\}$, where we assume N = 10. Dynamics within a country are

(2)

555

556 $dS_i/dt = \lambda_{S,i}X - (u + \tau_i)S$

 $X_i = 1 - S_i - R_i$

557
$$dR_i/dt = (1-c)\lambda_{R,i}X - uR$$

559

560 where we assume that treatment rates of subpopulations within a country 561 approximately follow a gamma distribution with shape parameter κ and mean 562 treatment rate τ . Accordingly, the rate of antibiotic consumption in subpopulation *i* is $\tau_i = \int_{Q_{\Gamma}\left(\frac{i}{\kappa} \mid \kappa\right)}^{Q_{\Gamma}\left(\frac{i}{\kappa} \mid \kappa\right)} t P_{\Gamma}(t \mid \kappa) dt, \text{ where } Q_{\Gamma}(q \mid \kappa) \text{ is the quantile } q \text{ of the gamma distribution}$ 563 564 with shape κ and $P_{\Gamma}(t|\kappa)$ is the probability density at *t* of the same gamma distribution. 565 At the scale of individuals, antibiotic consumption is highly variable, with some people 566 taking no antibiotics in a given year and others taking many courses of antibiotics (73); 567 at regional scales, antibiotic consumption shows less extreme variability (74) and 568 approaches a normal distribution. We use a gamma distribution to model variation in 569 treatment rates among subpopulations because it can capture patterns at either of these 570 scales, or scales in between. 571

572 The "Pathogen diversity" model extends the simple model (eq. 1) by structuring the 573 pathogen population into *D* different "D-types" (we assume D = 25), each with a 574 different natural clearance rate, where each type is kept circulating by diversifying 575 selection acting on D-type (*38*). Dynamics within a country are 576

577
$$dS_d/dt = q_d \lambda_{S,d} X - (u_d + \tau) S_d$$

578
$$\frac{dR_d}{dt} = q_d(1-c)\lambda_{R,d}X - u_dR_d$$

579
$$X = 1 - \Sigma_d (S_d + R_d)$$

where q_d is the strength of diversifying selection for D-type $d \in \{1, 2, ..., D\}$ and u_d is 581 582 the clearance rate for D-type d. We follow Lehtinen et al. (38) in defining $q_d =$

(3)

583
$$(1 - \frac{S_d + R_d}{\sum_{j=1}^D (S_j + R_j)} + \frac{1}{D})^a$$
 and $u_d = u \left(1 + \delta \left(2 \frac{d-1}{D-1} - 1 \right) \right)$, where *a* is the power of

584 diversifying selection and δ is the range of clearance rates. In a given country, the total 585 carriage of the type-*d* sensitive strain is S_d and the total carriage of the type-*d* resistant 586 strain is *R*_d.

587

588 Finally, the within-host competition models (26) allow hosts to carry a mix of both 589 strains. Hosts can carry the sensitive strain with a small complement of the resistant 590 strain (S_R) or the resistant strain with a small complement of the sensitive strain (R_S) . 591 Dynamics within a country are 592

593 $dS/dt = \lambda_S X - (u + \tau)S - k(1 - c)\lambda_R S + b_0 S_R$

594
$$dS_R/dt = k(1-c)\lambda_R S - (u+\tau)S_R + bR_S - b_0S_R$$

595 $dR_S/dt = k\lambda_S R - (u + \tau)R_S - bR_S$

596
$$dR/dt = (1-c)\lambda_R X - uR - k\lambda_S R + \tau(S_R + R_S)$$

597
$$X = 1 - S - R - S_R - R_S,$$
(4)

598

599 where *k* is the rate of co-colonisation relative to primary colonisation, *b* is the within-600 host growth benefit of sensitivity (i.e. the rate of the $R_S \rightarrow S_R$ transition), and b_0 is the 601 rate of the $S_R \rightarrow S$ transition. We follow Davies *et al.* (26) in setting $b_0 = 4b$. In a given country, the total carriage of the sensitive strain is $S + S_R$ and the total carriage of the 602 603 resistant strain is $R + R_s$. "Treatment competition" assumes the cost of resistance is 604 incurred by reduced transmission potential (b = 0 and c > 0), while "Growth 605 competition" assumes that the cost of resistance is incurred through decreased within-606 host growth (b > 0 and c = 0). 607

In equations 1, 3 and 4, the force of infection of a particular strain A in country m is $\lambda_A =$ 608 $\beta(fA_{tot|m} + (1-f)\sum_{\ell=1}^{M} h_{\ell}A_{tot|\ell})$, where β is the transmission rate, f is the between-609

country assortativity, h_{ℓ} is the relative population size of country *m* (such that $\sum_{\ell} h_{\ell} =$ 610 1), and $A_{tot|\ell}$ is the total carriage of strain A in country ℓ . The probability with which 611 individuals contact an individual from another country, 1 - f, captures those contacts 612 613 made with individuals from another country in either one's home country or a foreign country. In equation 2, the force of infection of a particular strain A in subpopulation *i* of 614 country *m* is $\lambda_{A,i} = \beta \left(f \left(g A_{tot|m,i} + (1-g) \sum_{j=1}^{N} \frac{1}{N} A_{tot|m,j} \right) + (1-g) \left(f \left(g A_{tot|m,i} + (1-g) \sum_{j=1}^{N} \frac{1}{N} A_{tot|m,j} \right) \right) \right)$ 615 $f \sum_{\ell=1}^{M} \sum_{j=1}^{N} \frac{h_{\ell}}{N} A_{\text{tot}|\ell,j}$, where g is the within-country assortativity and $A_{\text{tot}|\ell,j}$ is the 616 617 total carriage of strain A in subpopulation *j* of country ℓ . 618

619 Data and model fitting. We extracted community penicillin consumption and penicillin 620 non-susceptibility in *S. pneumoniae* invasive isolates from databases made available by 621 the ECDC (13, 28). We assume that community penicillin consumption drives penicillin 622 resistance, that antibiotic consumption is independent of whether an individual is 623 colonised by pneumococcus, and that resistance among invasive bacterial isolates is representative of resistance among circulating strains more broadly. Countries report 624 625 community penicillin consumption in defined daily doses (DDD) per thousand 626 individuals per day. To transform this bulk consumption rate into the rate at which 627 individuals undertake a course of antibiotic therapy, we analysed prescribing data from 628 eight European countries, estimating that, on average, 5 DDD in the population at large 629 correspond to one treatment course for a child under 5 years of age. This conversion 630 rate varies between countries (Table S3), but since the data are incomplete (8 of 27 631 countries) we have not explicitly accounted for this variability in our main model fitting 632 results.

633

634 Our model framework tracks carriage of *S. pneumoniae* among children aged 0–5 years, 635 the age group driving both transmission and disease. In European countries, we assume 636 that the prevalence of pneumococcal carriage in under-5s is 50% (11, 42) and the 637 average duration of carriage is 47 days (47, 48). We calculate the average incidence of S. 638 *pneumoniae*-caused severe pneumonia requiring hospitalisation as 610 per million 639 children under 5 per year (58) across the European countries in our data set. See Tables 640 S4, S5, and S6 for details of calculations relating to pneumococcal carriage duration and 641 disease incidence.

643 We use Bayesian inference via differential evolution Markov chain Monte Carlo (75) to

644 identify model parameters that are consistent with empirical data while accounting for

- 645 uncertainty in those estimates. Country *m* has antibiotic treatment rate τ_m and reports
- 646 r_m of n_m isolates are resistant. Over all *M* countries, these data are denoted $\tau =$

647 $(\tau_1, \tau_2, ..., \tau_M), r = (r_1, r_2, ..., r_M), \text{ and } n = (n_1, n_2, ..., n_M), \text{ respectively. The probability of}$ 648 a given set of model parameters θ is then

649

$$P(\theta|\tau,r,n) \propto P(\tau,r,n|\theta)P(\theta),$$

650

651 where $P(\theta)$ is the prior probability of parameters θ and

652
$$P(\tau, r, n|\theta) = C(Y = Y(\tau|\theta)) \prod_{m=1}^{M} R(r = r_m, n = n_m, \rho = \rho(\tau_m|\theta))^{N_m/\overline{N}}$$

653

654 is the likelihood of data τ , r, n given model parameters θ . Above, $Y(\theta)$ is the average 655 model-predicted prevalence of carriage across all countries and $\rho(\tau_m | \theta)$ is the modelpredicted resistance prevalence for country m. C(Y) is the credibility of prevalence of 656 657 carriage Y and $R(r,n,\rho)$ is the credibility of r out of n isolates being resistant when the 658 model-predicted resistance prevalence is ρ . For C(Y), we use a normal distribution with 659 mean 0.5 and standard deviation 0.002. For $R(r,n,\rho)$, we use $R(r,n,\rho) =$ $\int_{0}^{1} T(x|\mu = \rho, \sigma = \sigma(\theta)) {n \choose r} x^{r} (1-x)^{n-r} dx$, a binomial distribution where the 660 probability of success is modelled as a [0,1]-truncated normal distribution centred on ρ 661 662 and with standard deviation σ . The parameter σ captures the unexplained betweencountry variation in resistance frequency. Here, $T(x|\mu, \sigma) = \frac{\varphi(x|\mu, \sigma)}{(\Phi(1|\mu, \sigma) - \Phi(0|\mu, \sigma))}$, where 663 $\varphi(\mu,\sigma) = \frac{1}{\sqrt{2\pi\sigma^2}} exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$ is the untruncated normal PDF and $\Phi(\mu,\sigma) = \frac{1}{2}(1+\omega)$ 664 $erf\left(\frac{x-\mu}{\sigma\sqrt{2}}\right)$) is the untruncated normal cumulative distribution function. Finally, N_m is the 665 population size of country *m* and \overline{N} is the average population size across all countries; 666 the exponent N_m/\overline{N} allows us to weight the importance of each country by its 667 668 population size, which allows a closer fit with the overall resistance prevalence across 669 all countries. 670

- 671 As prior distributions for parameter inference, we adopt $c \sim \text{Beta}(\alpha = 1.5, \beta = 8.5)$, $b \sim \text{Gamma}(\kappa = 2, \theta = 0.5), \beta \sim \text{Gamma}(\kappa = 5, \theta = 0.35), g \sim \text{Beta}(\alpha = 10, \beta = 1.5),$ 672 $\kappa \sim \text{Gamma}(\kappa = 4, \theta = 2), a \sim \text{Gamma}(\kappa = 2, \theta = 5), \delta \sim \text{Beta}(\alpha = 20, \beta = 25), \text{ and}$ 673 674 $k \sim \text{Normal}(\mu = 1, \sigma = 0.5)$. Priors for *c*, *b*, and β were chosen to be vague since these 675 parameters are heavily constrained by the data we fit our models to. Priors for g, κ , a, δ , 676 and k were chosen to keep parameters within biologically plausible ranges. Table S7 provides more detail on the choice of priors, and Fig. S4 shows which parameters are 677 678 most strongly constrained by these prior beliefs.
- 679

680 We set the unexplained between-country variation in resistance prevalence σ to 0.06 681 across all models based on a preliminary round of model fitting with σ as a free 682 parameter. We set the between-country assortativity *f* to 0.985 (*i.e.*, 1.5% of contacts 683 occur with individuals from a different country) based on rates of travel for EU 684 residents. Specifically, using Eurostat database tour_dem_tnw (76) we estimated that 685 the average EU resident spent 1.5% of their nights abroad in 2007; this overestimates 686 mixing because children under 5 travel less than the average person, but 687 underestimates mixing because it does not account for contacts made with visitors to an 688 individual's country of residence and because children may contract pneumococcal 689 carriage from adults who travel, and so we kept the value of 1.5%. See Table S8 for 690 MCMC diagnostics.

691

692 To match model predictions to a high-burden setting, we increase the duration of

693 carriage to 71.4 days; increase the transmission rate by a factor of 3.49 (Treatment

diversity), 3.62 (Pathogen diversity), 3.61 (Treatment competition), or 3.20 (Growth

695 competition), so that carriage prevalence reaches 90.0%; and increase the antibiotic

696 consumption rate to 1.670, 1.458, 1.138, or 5.887 courses per person per year,

697 respectively, so that resistance prevalence reaches 81.4%.

698

699 *Interventions*. Interventions have the following impact on model parameters: for the

acquisition-blocking vaccine, the transmission rate becomes $\beta' = (1-\varepsilon_a)\beta$; for the

701 clearance-accelerating vaccine, the clearance rate becomes $u' = u/(1-\varepsilon_c)$; and under

antibiotic stewardship, the average treatment rate in each country *m* becomes $\tau_m' = \tau_m$

703 (1-ε_s).

705 *Capturing additional between-country variation in resistance frequency.* We begin by 706 finding the maximum *a posteriori* model fits according to the likelihood and prior 707 distributions for each of the four models of resistance evolution. This identifies the 708 following parameter values for each model. "Treatment diversity": $\beta = 1.41$, c = 0.124, g709 = 0.976, and κ = 2.22. "Pathogen diversity": β = 1.33, *c* = 0.191, *a* = 10.8, and δ = 0.608. 710 "Treatment competition": $\beta = 1.42$, c = 0.191, and k = 1.64. "Growth competition": $\beta = 1.42$ 711 1.39, b = 0.195, and k = 1.61. Then, we perform maximum *a posteriori* model fits for each 712 potentially-varying parameter under each model, allowing the varying parameter to 713 take on a different value for each country and fixing other parameters at their maximum 714 *a posteriori* values as determined in the previous step, or at specific assumed values for 715 u = 0.65, f = 0.985, and z = 5. For the second step, we use a modified likelihood function 716

717
$$P(\tau, r, n|\theta) = C(Y = Y(\tau|\theta)) \prod_{m=1}^{M} \phi\left(\mu = \frac{r_m + 1}{n_m + 2}, \sigma = 0.001 \middle| x = \rho(\tau_m|\theta)\right)^{N_m/\overline{N}},$$

718

719 where $\phi(\mu, \sigma | x)$ is the normal probability density function. This modified likelihood 720 function ensures that the model-predicted resistance frequency for each country is 721 matched as closely as possible to the maximum-likelihood resistance prevalence $\frac{r_m+1}{n_m+2}$ 722 (*i.e.*, assuming a uniform prior on resistance frequency) for each country *m*, so that 723 model fits are comparable across different varying parameters. We use the Nelder-Mead 724 algorithm to maximize the posterior probability in both steps.

725

Figs. S5–S8 show maximum *a posteriori* fits when allowing an additional parameter to
vary freely between countries, along with the parameter values identified by model
fitting. Figs. S9–S12 show the impact of vaccination, focusing on those parameters for
which model fitting was able to capture the observed variability in resistance frequency
between countries (*i.e.*, those parameters plotted to the left of the dashed line in Fig. 6b
of the main text). Figs. S13–S16 show the impact of vaccination for the remaining
parameters.

733	List of Supplementary Materials
734	
735	Fig. S1. Patterns of penicillin non-susceptibility across European countries, 2005–2017.
736	Fig. S2. Impact of key parameters upon potential for coexistence in each model.
737	Fig. S3. Carriage and resistance of D-types in "Pathogen diversity" model.
738	Fig. S4. Comparison of inferred model posteriors and conditional posterior maxima for
739	each parameter.
740	Fig. S5. Varying-parameter fits for the "Treatment diversity" model.
741	Fig. S6. Varying-parameter fits for the "Pathogen diversity" model.
742	Fig. S7. Varying-parameter fits for the "Treatment competition" model.
743	Fig. S8. Varying-parameter fits for the "Growth competition" model.
744	Fig. S9. Vaccine impact for the "Treatment diversity" model, varying parameters <i>c</i> and <i>z</i> .
745	Fig. S10. Vaccine impact for the "Pathogen diversity" model, varying parameters c , δ ,
746	and <i>z</i> .
747	Fig. S11. Vaccine impact for the "Treatment competition" model, varying parameters β ,
748	<i>c</i> , <i>u</i> , <i>k</i> , and <i>z</i> .
749	Fig. S12. Vaccine impact for the "Growth competition" model, varying parameters b and
750	Ζ.
751	Fig. S13. Vaccine impact for the "Treatment diversity" model, varying other parameters.
752	Fig. S14. Vaccine impact for the "Pathogen diversity" model, varying other parameters.
753	Fig. S15. Vaccine impact for the "Treatment competition" model, varying other
754	parameters.
755	Fig. S16. Vaccine impact for the "Growth competition" model, varying other parameters.
756	Table S1. Literature review.
757	Table S2. Summary of model parameters.
758	Table S3. Penicillin consumption.
759	Table S4. Carriage duration (Europe).
760	Table S5. Pneumococcal morbidity.
761	Table S6. Carriage duration (Kilifi).
762	Table S7. Priors for model fitting.
763	Table S8. MCMC diagnostics.
764	

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- 1025

1026	Acknowledgements
1027	
1028	Funding: N.G.D., M.J. and K.E.A. were funded by the National Institute for Health
1029	Research Health Protection Research Unit in Immunisation at the London School of
1030	Hygiene and Tropical Medicine in partnership with Public Health England. The views
1031	expressed are those of the authors and not necessarily those of the NHS, National
1032	Institute for Health Research, Department of Health or Public Health England. S.F. was
1033	supported by a Sir Henry Dale Fellowship jointly funded by the Wellcome Trust and
1034	Royal Society (grant number 208812/Z/17/Z).
1035	
1036	Author contributions: All authors designed the study. N.G.D. led the analysis with input
1037	from K.E.A., M.J., and S.F. The paper was written by N.G.D. and K.E.A. with input from M.J.
1038	and S.F.
1039	
1040	Competing interests: The authors declare no competing interests.
1041	
1042	Code availability
1043	
1044	C++ code used for model comparison is available at

1045 <u>https://github.com/nicholasdavies/amr-competition-diversity</u>.

1046	Figure captions
1047	
1048	Fig. 1. Four models of resistance evolution. X hosts are uncolonised, S hosts are
1049	colonised with the sensitive strain and R hosts are colonised with the resistant strain.
1050	Force of infection terms $\lambda_{\!A}$ are equal to the person-to-person contact rate β times the
1051	probability that a contacted individual carries strain <i>A</i> ; <i>c</i> is the transmission cost of
1052	resistance; u is the natural clearance rate; and $ au$ is the rate of antibiotic treatment. (a)
1053	"Treatment diversity": each country is split into subpopulations varying in treatment
1054	rate $ au_i$, with treatment rates drawn from a gamma distribution with shape к. Within a
1055	country, individuals assort with their own subpopulation with probability g ; this
1056	assortative mixing among treatment-varying subpopulations allows coexistence
1057	between sensitive and resistant strains. (b) "Pathogen diversity": the pathogen comes in
1058	multiple subtypes maintained by diversifying selection, each with its own natural
1059	carriage duration u_d^{-1} . Diversifying selection is stronger (<i>i.e.</i> , more equalizing) as a
1060	increases, while carriage durations span a greater range as δ increases. Only those
1061	subtypes whose carriage duration exceeds a critical threshold (dashed line) are selected
1062	for resistance, so that overall, both sensitive and resistant strains can circulate. (c)
1063	"Treatment competition": singly-colonised hosts can acquire a small amount of another
1064	strain at relative rate k (host states S_R and R_S). Dually-colonized hosts only transmit the
1065	dominant strain, so there is within-host competition between co-colonising strains.
1066	Population-level coexistence is maintained by treatment-mediated within-host
1067	competition. (d) "Growth competition": as in panel c, but the transmission cost of
1068	resistance is removed and sensitive strains now outgrow resistant strains within co-
1069	colonised hosts at rate <i>b</i> . Coexistence is maintained by both treatment-mediated and
1070	growth-mediated within-host competition. Panels a–d illustrate alternative model
1071	dynamics for a single country; our full model framework tracks dynamics for 27
1072	European countries simultaneously, which themselves mix with assortativity <i>f</i> .
1073	
1074	Fig 2. Four models reproduce patterns of resistance in <i>S. pneumoniae</i> in Europe.
1075	(a) Model fits with associated WAIC (± standard error). Points and vertical lines show
1076	the mean and 95% highest density intervals (HDIs) for the reported proportion of

1077 invasive *S. pneumoniae* isolates that are resistant to penicillin plotted against the

1078 penicillin consumption rate in under-5s. Ribbons show the 50% and 95% HDIs for

- resistance prevalence from each fitted model. (b) The top row shows estimated
 posterior distributions for the free parameters in each model; the bottom row shows
- 1081 model outputs associated with these parameters to aid interpretation.
- 1082

Fig. 3. Impact of interventions. Impact of vaccine and treatment interventions on (a)
carriage prevalence, (b) resistance frequency, and (c) resistant carriage (mean and
95% HDI). (d) Illustration of the strongest forces selecting for greater or lesser
resistance across models.

1087

1088 Fig. 4. Policy considerations. (a) Median equivalent reduction in prescribing across 1089 four models of resistance evolution, in terms of vaccine efficacy at reducing the 1090 prevalence of resistant pneumococcal carriage. This demonstrates the vaccine efficacy 1091 required to achieve a similar decrease in resistant carriage to a given reduction in 1092 antibiotic prescription rates. The impact on overall pneumococcal carriage is not 1093 considered here. The shaded bar shows an 8.8-23.1% reduction in prescriptions, an 1094 estimate of the percentage of prescriptions which are clinically inappropriate in the UK 1095 (77). The dashed line shows a 15% reduction in prescriptions, which has recently been 1096 announced as a target by the UK government (52). (b) The full impact of vaccination, 1097 illustrated here with 30% vaccine efficacy, can take 5-20 years to play out (mean and 1098 95% HDI). (c) Per-country impact of vaccination at 30% efficacy. Countries reporting to 1099 ECDC are ordered from lowest (NL) to highest (CY) reported rate of penicillin 1100 consumption. Diamonds show the estimated change in all pneumococcal pneumonia 1101 cases, while filled distributions show the change in resistant cases.

1102

Fig. 5. Vaccine impact in a high-burden setting. Adjusting fitted models to be
consistent with a high-burden setting yields different predictions for vaccine impact
(mean and 95% HDI), highlighting both increased challenges and greater opportunities
for resistance management via vaccination.

1107

1108 **Fig 6. Explaining additional between-country variation in resistance frequency.**

- 1109 Allowing model parameters to vary across countries captures additional between-
- 1110 country variation in resistance frequency not captured by variation in the treatment
- 1111 rate. For example, (a) allowing the transmission rate β to vary across countries can

- 1112 explain the variation in some but not all models (points and vertical lines show mean
- 1113 credible resistance frequency and 95% HDI for each country). **(b)** Depending upon
- 1114 which parameter is allowed to vary, models differ in how well they explain all additional
- 1115 between-country variation, with a clear separation (dashed line) between flexible and
- 1116 inflexible models. **(c)** Model-specific predictions for the impact of vaccination among
- 1117 those parameters that do fully capture the observed variation remain similar. Solid grey
- 1118 lines show "base" model; dashed lines correspond with colours in panel b.

Tables

Table 1. Mechanisms for maintaining coexistence

	Mechanism	Mode of action	Plausible mechanism for coexistence in <i>S. pneumoniae</i> ?	Consistent with empirical patterns? (Fig. 2)	
	Treatment diversity	Assortatively-mixing subpopulations differ in treatment rates (<i>25, 34–37</i>)	Ves Yes	~	Yes
	Pathogen diversity	Subtypes maintained by diversifying selection differ in propensity for resistance (38)	Ves Yes	~	Yes
	Treated class	Individuals currently in treatment maintain resistant strains (<i>25, 34, 39, 40</i>)	No: Only supports a small amount of coexistence (25)		N/A
IVERSITY	Within-host niches	Sensitive and resistant strains exploit separate niches within the host (<i>30</i> , <i>41</i>)	No: Resistant and sensitive strains are known to occupy the same niches (29)		N/A
IQ	Mutation pressure	Mutation-selection balance maintains intermediate resistance frequency (<i>30, 31,</i> <i>37</i>)	No: De novo acquisition of resistance in <i>S. pneumoniae</i> is not frequent enough (<i>25</i>)	-	N/A
	Prescription feedback	Doctors reduce prescribing of a drug as resistance to it increases (<i>37, 39</i>)	No: Does not explain how coexistence is maintained over a range of treatment rates		N/A
Z	Within-host competition: Treatment competition	Within-host competition creates frequency-dependent selection for resistance (25, 26, 32, 33, 40)	Ves Yes	~	Yes
COMPETITIC	Within-host competition: Growth competition	и	Ves Yes	~	Yes
	Superinfection	Superinfection creates frequency-dependent selection for resistance (30)	No: Requires resistant strain to transmit better than sensitive strain in absence of antibiotics	-	N/A

1124	Supplementary Materials for
1125	
1126	Stabilising effects of competition and diversity
1127	determine vaccine impact on antibiotic resistance evolution
1128	
1129	Nicholas G. Davies, Stefan Flasche, Mark Jit, Katherine E. Atkins







1132 Fig. S1. Patterns of penicillin non-susceptibility across European countries, 2005–

1133 **2017. (a)** The frequency of penicillin non-susceptibility in pneumococcal isolates (mean

and 95% HDI) increases with the primary care consumption of penicillins, ATC class

1135 J01C (least-squares linear regression $\beta = 0.0168$, F(1,344) = 148.5, $P < 2.2 \times 10^{-16}$).

1136 Data from Belgium after 2007 are excluded because of changes to the definition of

- 1137 penicillin non-susceptibility after this year. **(b)** The frequency of penicillin non-
- susceptibility has fluctuated in individual countries between 2005 and 2017, but the
- 1139 European population-weighted average (thick dashed line) has remained stable at
- roughly 12%. (c) We fit our models to the prevalence of penicillin non-susceptibility in
- 1141 2007. Despite the introduction of pneumococcal conjugate vaccines into many

- 1142 European national immunisation programmes starting in 2006, pneumococcal
- resistance appears to have been relatively stable over the time period 2005–2009:
- 1144 while mean penicillin non-susceptibility across European countries increased by 0.5%
- 1145 per year over this time period, and the slope of penicillin non-susceptibility on primary-
- 1146 care penicillin consumption decreased by 0.1%/DDD per year, neither of these changes
- 1147 are significant (least-squares linear regression: F(1,3) = 0.64, P = 0.48 and F(1,3) = 1.48,
- 1148 P = 0.31, respectively).
- 1149



1151 Fig. S2. Impact of key parameters upon the potential for coexistence in each

1152 **model. (a)** Potential for coexistence for each model, depending upon key parameters. In

1153 the top two panels, darker colours represent lower potential for coexistence while

1154 lighter colours represent higher potential for coexistence. **(b)** Schematic showing how

- 1155 to calculate the "potential for coexistence", an ad-hoc measurement of how much
- 1156 coexistence is exhibited by a given model under a given parameterisation. To find it, we
- 1157 set β = 1.4 months⁻¹, *u* = 0.7 months⁻¹, and set *g*, κ, δ, *a*, and *k* according to the values
- shown in the figure (panel a). We then numerically identify the value of *c* (or *b* for the
- 1159 "growth competition" model) which results in the equilibrium resistance frequency
- 1160 passing through exactly 30% for a single country in which the antibiotic consumption

- 1161 rate is equal to $\tau = 1.2$ courses per person per year (black dot; these values are chosen
- 1162 to approximately match observations for *S. pneumoniae* in European countries). The
- 1163 potential for coexistence is the probability that the equilibrium resistance frequency is
- above 0 for a random treatment rate between $\tau = 0$ and $\tau = 1.2$ (in other words, it is the
- 1165 proportion of the curve that "lifts off" from the x-axis; see figure). By this definition, a
- 1166 model showing no coexistence has potential 0, while a model showing the maximum
- amount of coexistence has potential 1.



- 1170 Fig. S3. Carriage and resistance of D-types in the "Pathogen diversity" model. This
- 1171 verifies that the D-types with the highest prevalence of carriage (averaged over all
- 1172 countries, above) also exhibit the highest resistance frequency (separated by country,
- 1173 below), and shows that at equilibrium, D-types can exhibit intermediate frequencies of
- 1174 resistance.



1176 Fig. S4. Comparison of inferred model posteriors and conditional posterior 1177 maxima for each parameter. The conditional posterior maximum (green bars) is 1178 obtained by fixing the parameter of interest at a specific value, then allowing the other 1179 parameters to assume their maximum *a posteriori* values through numerical 1180 optimization of the model fit. This is repeated for multiple values of the parameter of 1181 interest. The position on the x-axis of the thin green bars shows these fixed values for 1182 the parameter of interest (chosen to "sweep" across the parameter range), while the 1183 height of the thin green bars is relative to the maximum posterior probability conditional on that fixed value. When the conditional posterior maximum (green bars) 1184 does not align with the posteriors from model fitting (blue bars, same as Fig. 2b, main 1185 1186 text), this indicates that the prior distribution is strongly influencing the posterior 1187 distribution for a given parameter. This analysis shows that values inferred for *a* and δ under the "Pathogen diversity" model, and for *k* under the "Treatment competition" and 1188 "Growth competition" models, are strongly constrained by prior beliefs about plausible 1189 1190 ranges for those parameters, while other parameters are less constrained. 1191



- 1196 between countries. Parameters *c* and *z* can capture the additional variation in resistance
- 1197 frequency between countries.



- *posteriori* fits for the "Pathogen diversity" model allowing one parameter to vary
- 1202 between countries. Parameters c, δ , and z can capture the additional variation in
- 1203 resistance frequency between countries.



1206Fig. S7. Varying-parameter fits for the "Treatment competition" model. Maximum a1207posteriori fits for the "Treatment competition" model allowing one parameter to vary1208between countries. Parameters β , c, u, k, and z can capture the additional variation in1209is the experiment of t

1209 resistance frequency between countries.



Fig. S8. Varying-parameter fits for the "Growth competition" model. Maximum *a*

- *posteriori* fits for the "Growth competition" model allowing one parameter to vary
- 1214 between countries. Parameters *b* and *z* can capture the additional variation in
- 1215 resistance frequency between countries.

Treatment diversity



1216

1217

1218 Fig. S9. Vaccine impact for the "Treatment diversity" model, varying parameters *c*

- 1219 **and z.** Impact of vaccination under the "Treatment diversity" model, for those
- 1220 parameters able to capture the between-country variation in resistance frequency. The
- 1221 base model fit (thick grey solid line) is compared with the model fits in which
- 1222 parameters vary between countries (thin dashed lines).

Pathogen diversity



- 1223
- 1224

1225 Fig. S10. Vaccine impact for the "Pathogen diversity" model, varying parameters *c*,

- 1226 **δ, and z.** Impact of vaccination under the "Pathogen diversity" model, for those
- 1227 parameters able to capture the between-country variation in resistance frequency. The
- 1228 base model fit (thick grey solid line) is compared with the model fits in which
- 1229 parameters vary between countries (thin dashed lines).

Treatment competition



- 1230
- 1231

1232 Fig. S11. Vaccine impact for the "Treatment competition" model, varying

1233 **parameters** *β*, *c*, *u*, *k*, and *z*. Impact of vaccination under the "Treatment competition"

1234 model, for those parameters able to capture the between-country variation in resistance

- 1235 frequency. The base model fit (thick grey solid line) is compared with the model fits in
- 1236 which parameters vary between countries (thin dashed lines).

Growth competition



- 1237
- 1238

1239 Fig. S12. Vaccine impact for the "Growth competition" model, varying parameters

- 1240 *b* and *z*. Impact of vaccination under the "Growth competition" model, for those
- 1241 parameters able to capture the between-country variation in resistance frequency. The
- 1242 base model fit (thick grey solid line) is compared with the model fits in which
- 1243 parameters vary between countries (thin dashed lines).

Treatment diversity



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1245

1246 Fig. S13. Vaccine impact for the "Treatment diversity" model, varying other

1247 **parameters.** Impact of vaccination under the "Treatment diversity" model, for those

1248 parameters *not* able to capture the between-country variation in resistance frequency.

- 1249 The base model fit (thick grey solid line) is compared with the model fits in which
- 1250 parameters vary between countries (thin dashed lines).

Pathogen diversity



1251

1252

1253 Fig. S14. Vaccine impact for the "Pathogen diversity" model, varying other

- 1254 **parameters.** Impact of vaccination under the "Pathogen diversity" model, for those
- 1255 parameters *not* able to capture the between-country variation in resistance frequency.
- 1256 The base model fit (thick grey solid line) is compared with the model fits in which
- 1257 parameters vary between countries (thin dashed lines).

Treatment competition



1258

1259



1261 **parameters.** Impact of vaccination under the "Treatment competition" model, for those

1262 parameters *not* able to capture the between-country variation in resistance frequency.

- 1263 The base model fit (thick grey solid line) is compared with the model fits in which
- 1264 parameters vary between countries (thin dashed lines).

Growth competition



1265

1266

1267 Fig. S16. Vaccine impact for the "Growth competition" model, varying other

1268 **parameters.** Impact of vaccination under the "Growth competition" model, for those

1269 parameters *not* able to capture the between-country variation in resistance frequency.

1270 The base model fit (thick grey solid line) is compared with the model fits in which

- 1271 parameters vary between countries (thin dashed lines).
- 1272

1273	Tables S1–S7
1274	
1275	These tables can be found in an Excel spreadsheet accompanying the article.
1276	
1277	Table S1. Literature review — Details of the literature review used to identify
1278	mechanisms for maintaining coexistence between sensitive and resistant bacterial
1279	strains.
1280	
1281	Table S2. Summary of model parameters — Table describing model parameters and
1282	assumed values or prior distributions for model fitting.
1283	
1284	Table S3. Carriage duration — Calculation of mean pneumococcal carriage duration for
1285	children under 5 years old in European settings.
1286	
1287	Table S4. Penicillin consumption — Calculation of the mean number of defined daily
1288	doses of penicillin corresponding to a single treatment course for children under 5
1289	years old in European countries.
1290	
1291	<i>Table S5. Pneumococcal morbidity</i> — Calculation of the annual number of pneumococcal
1292	pneumonia cases in children under 5 in Europe and Kenya.
1293	
1294	Table S6. Carriage duration (Kilifi) — Calculation of mean pneumococcal carriage
1295	duration for children under 5 years old in Kilifi, Kenya.
1296	
1297	Table S7. MCMC diagnostics — Widely Applicable Information Criteria (WAIC), Leave-
1298	One-Out Information Criteria (LOOIC), effective posterior sample size and Gelman-
1299	Rubin diagnostics for Bayesian inference model fitting using Markov chain Monte Carlo.
1300 1301	