Within-host dynamics

Understanding within-host host-parasite interactions (focus on dynamics)

Lots of molecular biology, genetics (recognition mechanisms and effector mechanisms), won't deal with that now.

Interaction between different components of the immune system (modeled at different levels of detail/realism), parasite populations (maybe in multiple compartments?)

Longitudinal data (relatively rare), distributional data.

HIV dynamics under (ineffective) treatment

Bonhoeffer, Coffin, and Nowak (1997)

- Early HIV antivirals: relatively ineffective due to rapid mutation
- Large decline in virus loads (up to 300-fold decline in viral RNA in some patients)
- but no clearance
- within-host $R_0 \approx 50$

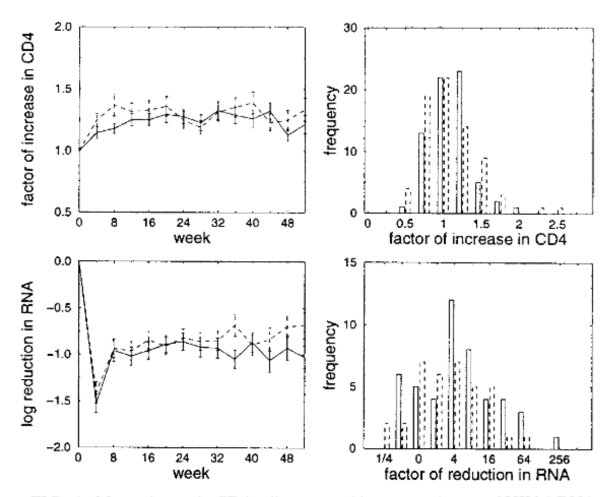
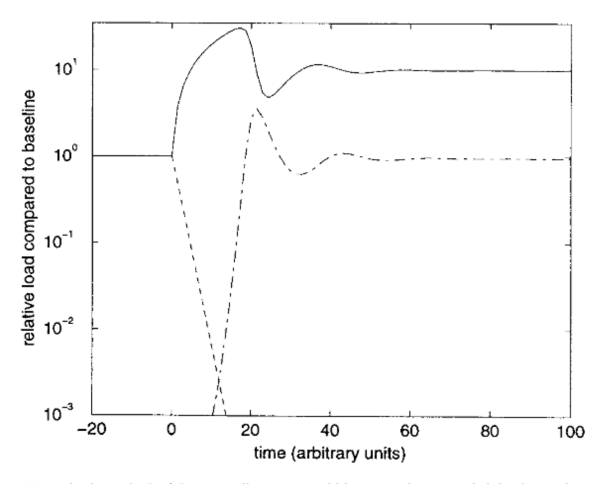


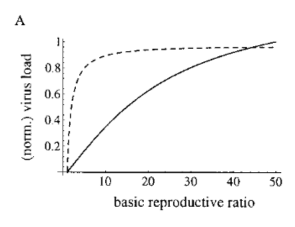
FIG. 1. Mean change in CD4 cell count and log mean change of HIV-1 RNA load compared to baseline in patients treated with a low-dose (dashed line) and high-dose (solid line) combination of lamivudine and zidovudine. In each treat-

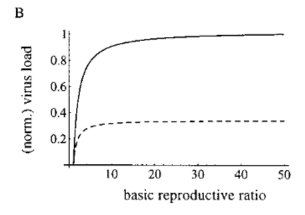


• "virus load paradox": if R_0 is initially 50, we would have to reduce it to slightly above 1 but never **below** 1 to see these results.

$$\frac{dC}{dt} = \lambda - \mu C - \beta CV$$
$$\frac{dV}{dt} = \beta CV - aV$$

- add a drug-resistant type to the model
- $\bullet\,$ add mutation (and back-mutation) to the model
- add immune responses $(dz/dt = kV \gamma z)$
- homeostasis of infectible cells (logistic growth)
- virus-induced killing of uninfected cells (e.g. gp120 shedding)
- differential effects of drug on different types
- distribution of infectibility



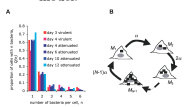


Within-host (and within-cell) dynamics of salmonella

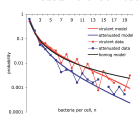
• intracellular bacterium

Brown et al. (2006)

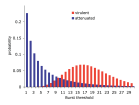
• model:



• assume that host cells are always available (infinite S)



• distribution: two categories, or a range of burst sizes?



- "constitutive" vs "stochastic" models
- density-dependence in growth and/or burst probability?
- extracellular killing (bactericidal) vs slowing/preventing intracellular growth (bacteriostatic)

our analysis predicts that the efficacy of common extracellular antibiotics can be enhanced by supplementation with antibiotics slowing intracellular bacterial division [bacteriostatic drugs]. This implies that both bacteriostatic and bactericidal drugs can potentiate the therapeutic efficacy of extracellular antibiotics.

References

Bonhoeffer, S, J M Coffin, and M A Nowak. 1997. "Human Immunodeficiency Virus Drug Therapy and Virus Load." Journal of Virology 71 (4): 3275–78. http://www.ncbi.nlm.nih.gov/pmc/articles/PMC191463/. Brown, Sam P, Stephen J Cornell, Mark Sheppard, Andrew J Grant, Duncan J Maskell, Bryan T Grenfell, and Pietro Mastroeni. 2006. "Intracellular Demography and the Dynamics of Salmonella Enterica Infections." PLoS Biol 4 (11): e349. https://doi.org/10.1371/journal.pbio.0040349.

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