Additional file 1: Text S1. Cluster age profiles

To check for the existence of a consistent age profile pattern of prevalence across clusters in the baseline data, we employ a simple model of age-dependent prevalence, $p_i(a)$, in cluster $i$,

$$p_i(a; m) = \bar{p}_i + m(a - \bar{a}_i)$$

where $m$ is the gradient of prevalence with age and $\bar{p}_i$ and $\bar{a}_i$ are cluster-specific intercept values. The value $\bar{p}_i$ is the overall prevalence in cluster $i$ from the data and

$$\bar{a}_i = \frac{1}{n_i} \sum_j a_{ij}$$

where $n_i$ is the sample size from cluster $i$ and $a_{ij}$ is the age of the $j^{th}$ individual in the $i^{th}$ sample. This definition has the advantage that average prevalence from the model across the age points in the data always matches that from the data itself. We calculate a log-likelihood for each cluster,

$$L_i(m) = \sum_{j \in S_i^+} \ln p_i(a_{ij}; m) + \sum_{j \in S_i^-} \ln(1 - p_i(a_{ij}; m))$$

where $S_i^+$ and $S_i^-$ are the sets of the ages of all egg-positive and egg-negative individuals in the $i^{th}$ sample, respectively. The total log-likelihood is

$$L(m) = \sum_i L_i(m).$$

Fig S1A shows the log-likelihood for the age profile prevalence gradient over all clusters, $L(m)$. The maximum likelihood estimator (MLE) for the gradient is $m = 0.0012$/year with a very small degree of uncertainty, implying a prevalence difference of approximately 6% across a 50 year age difference. This matches the gradient seen in the age profile of the data over all (Fig 1D in the main text). From the point of view of individual clusters, however, there is considerable variability. Fig S1B shows a histogram of the MLE values for gradients in individual clusters. Values cluster around the overall best fit gradient, but are clearly not significantly different from zero, which would imply no age structure. Additionally, the posterior distributions for cluster-level gradient estimates have standard deviations of around $\approx 0.0015$/year. These observations, together with the additional simplicity and fewer parameters, support the decision to use a transmission model without age structure.
Figure S1: a) Total log-likelihood for prevalence gradient, $m$ across all clusters (MLE: $m = 0.0012/\text{year}$). b) histogram of MLE prevalence gradients for the 119 clusters. Individual MLE values are subject to an uncertainty of $\approx 0.0015/\text{year}$.