Assessing the stability of polio eradication after the withdrawal of oral polio vaccine

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Abstract

The oral polio vaccine (OPV) contains live-attenuated polioviruses that induce immunity by causing low virulence infections in vaccine recipients and their close contacts. Widespread immunization with OPV has reduced the annual global burden of childhood paralysis by a factor of ten thousand or more and has driven wild poliovirus (WPV) to the brink of eradication. However, in instances that have so far been rare, OPV can paralyze vaccine recipients and generate vaccine-derived polio outbreaks. To complete polio eradication, OPV use should eventually cease, but doing so will leave a growing population fully susceptible to infection. If poliovirus is somehow re-introduced after OPV cessation, will it be likely to cause outbreaks? If the virus is from OPV, will the risk of vaccine-derived outbreaks always remain low? To answer these questions, we built a multiscale mathematical model of infection and transmission with data from clinical trials and field epidemiology studies. The within-host model describes the effects of vaccination and waning immunity on shedding and dose response. The transmission model captures how shedding and dose response interact with sanitation and person-to-person contact patterns to determine the force of infection in communities. Our results show that inactivated polio vaccine alone is sufficient to prevent outbreaks in low force of infection settings, and that OPV can be started and stopped safely in moderate force of infection settings. However, in high force of infection settings, the conditions that support vaccine-derived outbreaks have only been rare because population immunity has been high. With insufficient immunity, the Sabin strains from OPV will be nearly as capable of causing outbreaks as WPV. We conclude that while eradication may still be achieved with continued use of Sabin OPV, a new vaccine is needed to secure eradication for all time.
Author Summary

Oral polio vaccine (OPV) has played an essential role in the elimination of wild poliovirus (WPV). OPV contains transmissible viruses that can spread from person-to-person. When OPV transmission persists uninterrupted, vaccine-derived outbreaks occur. After OPV is no longer used in routine immunization, as with the cessation of type 2 OPV in 2016, population immunity will decline. A key question is how this affects the potential of OPV viruses to spread within and across communities. To address this, we examined the roles of immunity, sanitation, and social contact in limiting OPV transmission. Our results derive from an extensive review and synthesis of vaccine trial data and community epidemiological studies. Shedding, dose response, and transmission data are analyzed to systematically explain and model observations of WPV and OPV circulation. We show that in high force of infection settings, falling population immunity after OPV cessation will lead to conditions where OPV and WPV are similarly capable of causing outbreaks, and that this conclusion is compatible with the known safety of OPV prior to global cessation.

Abbreviations

WPV, wild poliovirus; OPV, oral polio vaccine; tOPV, trivalent OPV; mOPV, monovalent OPV; bOPV, bivalent type 1 and 3 OPV; cVDPV, circulating vaccine-derived poliovirus; IPV, inactivated polio vaccine; CID50, the culture infectious dose that induces a cytopathic effect in 50% of infected cell or tissue cultures; HID50, the dose (measured in CID50) that infects 50% of orally-exposed and immunologically-naive humans; MLE, maximum likelihood estimate; CI, confidence interval; UP, Uttar Pradesh.

Introduction

Wild polioviruses (WPV) have been eliminated from all but three countries [1,2] by mass vaccination with the oral polio vaccine (OPV). The annual burden of paralytic polio infections has been reduced ten-thousand-fold since the start of vaccination efforts [1]. OPV has been the preferred vaccine for polio eradication because it costs less, can be reliably delivered by volunteers without medical training, and is more effective against poliovirus infection, relative to the inactivated polio vaccine (IPV) [3,4]. Unique among current human vaccines, the live-attenuated Sabin poliovirus strains in OPV are transmissible. This transmissibility provides additional passive immunization that
enhances the effectiveness of OPV for generating herd immunity. However, the attenuation of Sabin OPV is unstable and so it can, in rare instances, cause paralytic poliomyelitis [5] and lead to outbreaks of circulating of vaccine-derived poliovirus (cVDPV) with virulence and transmissibility comparable to that of wild poliovirus (WPV) strains [6]. Thus, to complete the task of poliovirus eradication, vaccination with Sabin OPV must eventually cease [7].

The dual role of Sabin OPV as both a vaccine and a source of poliovirus is responsible for key uncertainties surrounding the ability of the Global Polio Eradication Initiative to achieve and sustain eradication. Since the widespread introduction of polio vaccination, polio outbreaks have taken place in regions of low immunity against infection surrounded by regions of high immunity [8], OPV campaigns implemented in outbreak response have been effective for interrupting transmission [3], and cVDPV outbreaks have been rare consequences of the hundreds of millions of OPV doses administered every year [9].

However, after vaccination with OPV is stopped, population immunity against infection will progressively decline. If polioviruses are reintroduced, whether due to accidental or deliberate release [10–12], or because of sustained yet undetected transmission [2,13–15], then large outbreaks may again occur. Outbreak control will require vaccination campaigns in affected countries and perhaps much more broadly. Since the withdrawal of type 2 OPV in April 2016, realizations of this scenario have occurred in Nigeria [16,17], the Democratic Republic of Congo, Pakistan, and Syria [18]. In response, millions of doses of monovalent type 2 OPV (mOPV2) have been delivered in these and neighboring countries [19], and none of the outbreaks have been declared interrupted at the time of writing.

In this paper, we explore the implications of the accumulated evidence about polio infection and transmission for the long-term stability of polio eradication. Will it be possible to interrupt all polio outbreaks without restarting widespread OPV vaccination, now and at all times in the future? Fundamental to this question is the ability of the Sabin polioviruses to circulate in low immunity populations. After OPV cessation, under what conditions will Sabin OPV remain the most effective tool for eliminating outbreaks without significant risks of causing more?

To address these questions, and building from primary literature and previous reviews and models [4,20–25], we developed a comprehensive synthesis of the evidence for how within-host immunity, viral infectivity, and transmission dynamics fit together to explain the epidemiology of poliovirus transmission. We built a within-host model that summarizes the effects of immunization on poliovirus shedding and susceptibility. We then incorporated the within-host dynamics into a
poliovirus transmission model using a household–community network framework, and we calibrated the model to field transmission studies. With the model, we explored how the average force of infection in exposed communities varies with immunity, sanitation, number of social contacts, and poliovirus type. We identified conditions required for the Sabin strains to remain indefinitely as highly effective vaccines with low risks of causing outbreaks, and conditions were they can be expected to transmit nearly as efficiently as WPV. Our results are discussed in the context of the established stability of OPV cessation in the developed world and the ongoing global Sabin 2 cessation.

Previous models have also explored the effects of OPV cessation on Sabin and WPV transmission [20,26,33]. Our work shares a similar level of scholarship on individual immunity and infection dynamics [20,24,34], but it differs in its emphasis of epidemiological data that can be measured in the field. The earlier work used compartmental models that assume well-mixed, large populations [20,26,27,33] with either unrealistic sterilizing immunity [20,26,33] or substantial complexity that precluded rigorous calibration to data [29]. In contrast, our model describes person-to-person transmission with realistic effects of immunity. It is based on correlates of transmission that are often measured in epidemiological studies, permits full uncertainty propagation, and is designed to draw inferences about community transmission from contact-tracing data, both historical [35,36] and modern [37,38].

Methods

Overview. To integrate knowledge of within-host immunity, shedding, and acquisition with between-host transmission, we built a multi-scale mathematical model. We first performed a quantitative literature review of clinical trial data to determine the impact of polio vaccination schedules containing OPV and/or IPV on poliovirus shedding after challenge with OPV. This resulted in a statistical immune correlate—the OPV-equivalent antibody titer—which was used to model the associations between polio vaccination and shedding duration, concentration of virus in stool, and oral dose response. Second, we reviewed in detail three historical transmission studies to parameterize a model of poliovirus transmission within households and between close extra-familial contacts. We then extended the person-to-person model by defining the local reproduction number—a threshold parameter that summarizes the potential for epidemic transmission within a homogeneous community. The model was implemented in Matlab 2015b (The Mathworks, Natick,
MA) and is available at famulare.github.io/cessationStability/. For all model parameters, see Supplement.

We made simplifying assumptions while developing the model, all of which are revisited in the Results and Discussion. First, we did not include the oral-oral transmission route. The studies known to us show that oral shedding occurs for shorter durations than fecal shedding and most often in individuals with low immunity [39–42], and this route is likely important in high sanitation settings [34,40,43]. Second, we ignored fascinating questions about the effects of genetic evolution on Sabin strain transmission, and so our Sabin transmission parameters are most applicable in the first few weeks after OPV vaccination [44–46]. Third, our model focuses on transmission from children to family members and extrafamilial contacts, and it ignores other person-to-person interactions and possible environmental transmission routes, all of which influence the absolute probability and severity of outbreaks [47–51]. Fourth, since paralysis has no direct influence on transmission, we did not model the impact of vaccination on paralysis (see Vidor et al [52] for a review).

All model features that describe the fraction of subjects shedding after live poliovirus exposure were fit by maximum likelihood assuming binomial sampling. Models for positive-definite quantities (concentration of poliovirus in stool, antibody titer) were estimated by ordinary least squares on log(quantity), and 95% confidence intervals assume log-normality. 95% confidence intervals were estimated by parametric bootstrap with 1000 replicates. To estimate bootstrap confidence intervals of parameters that are conditionally-dependent on previously estimated parameters, we propagated uncertainty by resampling known parameters from the 95% confidence intervals prior to resampling the data and re-estimating the parameters under investigation. Model equations and more detailed discussions of design decisions and calibration results can be found in Supplement. Differences in comparable quantities are considered statistically significant at $\alpha = 0.05$.

**Within-host model**

Our within-host model describes shedding from and susceptibility to poliovirus infection. In the model, the ability of an infected individual to transmit polio depends on the duration of shedding and the concentration of poliovirus in their stool. Susceptibility to infection depends on the dose response relationship for the probability that poliovirus ingested orally results in an infection, as detected by subsequent fecal shedding. Shedding duration, concentration, and dose response all depend on pre-exposure immunity and the poliovirus source, vaccine or wild.
Immunity in our model is represented by the OPV-equivalent antibody titer (denoted \( N_{\text{Ab}} \))—a statistical correlate of immunity that is inferred from measurements of shedding duration and/or dose response (first introduced by Behrend et al \cite{25} and called “mucosal immunity” therein). Previous reviews have demonstrated that when immunity is due to prior OPV immunization or natural WPV infection, homotypic (of the same serotype) serum neutralizing antibody titers (measured as the geometric mean reciprocal dilution of serum that is able to neutralize 100 CID50 of poliovirus) are predictive of fecal shedding and susceptibility \cite{25,53}. However, serum antibody titers induced by IPV alone, and heterotypic titers against type 2 from bivalent type 1 and 3 OPV (bOPV), are not predictive of shedding and susceptibility \cite{25,54,55}. The OPV-equivalent antibody titer describes the impacts of vaccination histories containing IPV or bOPV on shedding and susceptibility in terms of equivalent serum antibody titers from homotypic OPV vaccination in children. This model is agnostic about the biophysical mechanisms of immunity that prevent fecal shedding and is not intended to represent IgA concentration or other direct correlates of mucosal immunity \cite{56}. Following the results of Behrend et al \cite{25}, we assumed that the typical immunologically-naive individual with no history of poliovirus exposure (“unvaccinated”) and no measurable humoral immunity (“seronegative”) has an OPV-equivalent antibody titer of \( N_{\text{Ab}} = 1 \) by definition, that the maximum typical OPV-equivalent titer is \( N_{\text{Ab}} = 2048 (= 2^{11}) \), and that homotypic antibody titers for each poliovirus serotype are independent.

The studies used to calibrate the within-host model span many countries, years, and types of immunization history. All included studies describe the fraction of subjects positive for poliovirus in stool after OPV challenge or WPV exposure as equal to the number of subjects shedding divided by the number tested at each time point. In many cases, the data were digitized from published figures that do not report variation in the number of samples for each time point, and so our sample sizes at each time point are often approximate. A summary of all included studies, with details about which studies contributed to which components of the model, and reasons for study exclusion appears in Supplement \cite{36,42,54,55,57–73}, and the data are in S1 Structured Code and Data.

**Shedding duration.** In Fig 1 we summarize available data for and our model of the impact of different immunization histories on shedding duration \cite{36,42,54,55,57,58,60,63,65,68,70,72}. Details about data aggregation and our log-normal survival model for shedding duration can be found in Supplement, including descriptions of the trial populations and the equation used to represent the fraction of people shedding over time (eq. (S1)). Subjects from all study arms included...
in this section were five years of age or younger. Note that waning and setting-specific variations in OPV efficacy were not considered at this stage of model building.

Our shedding duration model summarizes the following observations. In immunologically-naive individuals, there were no statistically significant differences by serotype in shedding duration after OPV challenge. The maximum likelihood estimate (MLE) of the median shedding duration for immunologically-naive individuals shedding any Sabin strain is 30.3 (23.6, 38.6) days, significantly shorter than the median shedding duration for WPV, 43.0 (35.7, 51.7) days. The median shedding duration associated with maximum OPV-equivalent antibody titer ($N_{Ab} = 2048$) is 6 (4, 10) days for the Sabin strains and is modeled to be 8 (6, 13) days for WPV. Repeated vaccination with IPV alone shows no cumulative effect on shedding duration and with study-to-study variation showing little or no effect overall. bOPV produces a decrease in shedding duration against heterotypic Sabin 2 challenge, and this effect may be weakly enhanced by IPV after bOPV but not IPV before. Repeated immunization with trivalent OPV (tOPV) has a cumulative effect on shedding duration. tOPV has weaker effects on shedding duration for type 3 than types 1 and 2, reflecting known efficacy differences \[74\]. At the level of aggregation examined here, limited data suggest mOPV is comparable to tOPV. The transformation from median duration (Fig 1B) to titer (Fig 1C) serves as the definition of the OPV-equivalent antibody titer in our model—short post-challenge shedding duration implies high pre-challenge OPV-equivalent immunity.

**Concentration of poliovirus in stool.** In Fig 2 we show available data for and our model of the concentration of poliovirus in stool while shedding after OPV challenge \[54, 55, 57, 59, 68, 69\]. The included studies all reported concentration as the geometric mean 50% culture infectious dose per gram of stool (CID50/g) averaged across all subjects positive for poliovirus at each time point, and individual-level variation was generally not reported. Ages at OPV challenge ranged from 6 months to 65 years or more. The majority of trial arms challenged subjects with mOPV2 (mOPV1, $n = 5$; mOPV2, $n = 11$; mOPV3, $n = 5$). Data exploration revealed no systematic differences in concentration by serotype. We are not aware of similar longitudinal data for WPV shedding. OPV-equivalent antibody titers were estimated from the corresponding shedding duration data for each trial arm (see Supplement), and trial arms considered immunologically-naive reported no history of live poliovirus exposure, contained confirmed seronegative subjects, or had OPV-equivalent antibody titers consistent with $N_{Ab} = 1$. 


Figure 1. Effect of vaccination on shedding duration after OPV challenge and pre-challenge immunity. Labels describe vaccines received and number of doses (i.e. bOPVx2 & IPVx1: two doses of bOPV and one dose of IPV). (A) Shedding duration after OPV challenge: shedding duration survival curves from aggregated trial data (solid lines) and maximum likelihood model fit (dashed) for each immunization schedule and poliovirus type. (Infection with Sabin 1, blue; Sabin 2, red; Sabin 3, orange; WPV, black). (B) Median shedding durations after infection due to OPV challenge or WPV transmission estimated from model fits in panel A and (C) corresponding pre-challenge homotypic OPV-equivalent antibody titers. (See also Supplement and an interactive visualization of shedding duration data is available at famulare.github.io/cessationStability/)
Figure 2. Concentration of poliovirus in stool: effects of age and immunity. (A) Mean concentration of polivirus in stool (CID50/g) vs. time after OPV challenge for immunologically-naive subjects (color by age at challenge). (B) Peak concentration depends on age (dot color by age at challenge, corresponding to data from panel A at one week post-challenge; green line, model MLE and 95% CI, eq. (S2)). (C) Mean concentration after mOPV2 challenge for subjects with various vaccination histories (dashed, model MLE; solid, trial data age-adjusted to 12 months using eq. (S3)). The concentration of poliovirus in stool depends on pre-challenge vaccination history. (D) The mean daily concentration (CID50/g/day) declines by one order of magnitude for every eight-fold increase in OPV-equivalent antibody titer (OPV-equivalent titers (color, MLE and 95% CI) for each trial arm shown in panel C; black, model (eq. (S4), MLE and 95% CI)). Interactive data visualization available at famulare.github.io/cessationStability/.
Our model of poliovirus concentration in stool summarizes the following observations. Poliovirus concentrations peak 5–8 days after acquiring infection and decline slowly thereafter (Fig 2A). Data from immunologically-naive subjects revealed an unexpected dependence of peak concentration with age (Fig 2B). Peak concentration declines by roughly two orders of magnitude over the first three years of life with an exponential aging constant of 12 (1, 45) months, consistent with major developmental milestones including the transition to solid food and immune system maturation [75,76], after which the limited data indicate stability of peak shedding concentration for life. The concentration of poliovirus in stool is correlated with vaccination history (Fig 2C) and decreases by roughly a factor of ten with each eight-fold increase in OPV-equivalent titer (Fig 2D). The shedding concentration model is described in eqs. (S2–S4).

Dose response. To inform our dose response model, we first examined studies of healthy children that measured the probability of fecal shedding after receiving oral droplets with doses ranging from $10^1$ to $10^6$ CID50, and for which pre-challenge immunization histories were known. Three studies challenged with Sabin 1 [42,62,64], none used Sabin 3 or WPV, and one unusual human-passage study challenged with Sabin 2 and type 2 poliovirus derived from Sabin 2 after five days of replication in vaccinated children [59]. There were no statistically significant differences between Sabin 1 and Sabin 2 across these trials, but statistical power at low doses is poor. We also included modern studies of vaccine doses ($10^5−6$ CID50) that provided information about the effects of heterotypic immunity against type 2 from bOPV [54,55] and IPV boosting on prior OPV immunization [73]. OPV-equivalent antibody titers were estimated from the corresponding shedding duration data for each trial arm.

Our dose response model summarizes the following observations (Fig 3A-B). The typical Sabin 1 dose required to infect 50% of immunologically-naive healthy children (the 50% human infectious dose, HID50) is 54 (26, 100) CID50, and the fraction shedding approaches one for doses greater than $10^4$ CID50. Immunity has similar effects on dose response as it does on shedding duration and concentration. IPV-only immunization reduces susceptibility to infection in some studies but not all, and the effect is at most comparable to that provided by heterotypic immunity against type 2 from immunization with bOPV. tOPV reduces susceptibility across all doses. Not addressed in previous sections on shedding is that IPV-boosting in subjects with prior OPV immunization is highly effective for reducing susceptibility—as is now well-known [73,77,78]. OPV-equivalent antibody titer has a monotonic relationship with dose response (Fig 3C). The data are consistent with an
immunity-dependent beta-Poisson dose response model [79] (Fig 3D and eq. (S5)).

**Figure 3. Dose response to oral OPV challenge.** (A) Fraction shedding after Sabin 1 oral challenge at different doses (color by trial arm, symbol by source study). (B) Fraction shedding after Sabin 2 oral challenge at different doses (color by trial arm, symbol by source study; data for doses \( \leq 10^3 \) CID50 are for human-passaged Sabin 2 isolated from stool five days after vaccination). (C) Fraction shedding at vaccine doses (\( 10^{5-6} \) CID50) decreases with increasing OPV-equivalent antibody titer. (Color and symbols as in panels A–B; black lines are model MLE and 95% CI using eq. (S5)). (D) Beta-Poisson model MLE and 95% CI. Three model scenarios shown correspond to immunologically-naive (\( N_{Ab} = 1 \), red), heterotypic bOPV and upper-bound IPV-only (\( N_{Ab} = 8 \), green), and typical tOPV or post-IPV-boosting (\( N_{Ab} = 256 \), blue). Data from panels A–B (symbols as above, colored by corresponding model scenario).

To inform our model of strain-specific differences in dose response, we examined two transmission studies from similar settings in the United States. The first study in Houston in 1960 [36] measured transmission among immunologically-naive close contacts of vaccinees for each of the Sabin strains, and another in Louisiana from 1953–55 [35] measured close contact transmission of WPV (combined across all serotypes); these studies and our transmission model are described in detail in a later Methods section. Under the assumptions that Sabin 1 is well-described by the OPV challenge model above and that sanitation and contact patterns are similar across the four trial arms, differences in transmission are attributable to the virus-specific differences in infectivity shown in Table 1.
Table 1. Infectivity by strain. MLE and 95% CI of the HID50—the oral dose that infects 50% of immunologically-naive people.

| strain | HID50     |
|--------|-----------|
| Sabin 1| 54 (26, 100) TCID50 |
| Sabin 2| 30 (15, 54) TCID50  |
| Sabin 3| 67 (34, 120) TCID50 |
| WPV    | 7 (2, 41) TCID50   |

Waning immunity. We built a composite picture of waning immunity against infection from analysis of OPV-equivalent antibody titers across studies. We considered data for individuals that were likely maximally immune after their last poliovirus exposure, either due to immunization with three or more doses of tOPV \[42,55,73\] or accumulated natural immunity through 15 years of age during the endemic era \[57,69\]. The included trial arms involved subjects from 6 months to 65+ years of age and with between 1 month and likely 45+ years from last immunizing event to OPV challenge (see Supplement for additional details).

Our waning model summarizes the following observations (Fig 4). Absent reinfection or vaccination, immunity declines over many years, possibly with increasing variation in adults. We modeled waning as a power-law decay \[80\] during the months since last immunization, \(N_{Ab}(t) \propto t^{-\lambda}\), with exponent \(\lambda = 0.87 (0.73, 1.02)\) (eq. (S6)). The limited relevant data after bOPV vaccination \[55\] are consistent with the hypothesis that heterotypic and homotypic immunity share waning dynamics (bOPV data only, \(\lambda = 0.52 (0, 1.2)\)).

Figure 4. Waning immunity against infection. OPV-equivalent antibody titer vs. time between last exposure and mOPV challenge (color by serotype and symbol by source of immunity). Power-law model of waning from maximum homotypic immunity (MLE and 95% CI, black lines) and heterotypic immunity against type 2 from bOPV (MLE and 95% CI assuming homotypic waning exponent, green lines).
Transmission model

Our model describes the effects of within-host dynamics on transmission among close social contacts. We assumed that transmission from infected person to recipient occurs by oral exposure to infected feces, where the amount of poliovirus transmitted per exposure is determined by the shedding duration and concentration models, and recipient susceptibility is determined by the dose response model.

**Person-to-person transmission.** The population structure of the model is based on the essential transmission network motif examined by the field transmission studies (Fig 5A): infected young children transmit to family members in their household, who transmit to their close social contacts outside the household. For each of the three individuals along the transmission chain, the person-to-person model calculates daily incidence (the probability of becoming infected each day), prevalence (the probability of shedding poliovirus in stool each day), and concentration of poliovirus shed (CID50 per gram of stool).

![Network motifs of poliovirus transmission](image)

**Figure 5.** Network motifs of poliovirus transmission. (A) The essential motif of poliovirus transmission is index child to family member to extrafamilial contact (dot color gives subject type, line color describes relationship). (B) The local reproduction number describes the expected number of extrafamilial households infected by an index child based on immunity, sanitation, and the number of extrafamilial contacts. The house-to-house transmission motif represents this first generation of local transmission (color as in panel A; gray boxes denote households; dashed lines indicate relationships beyond the first generation). (C) Our definition of the local reproduction represents the expected number of extrafamilial contacts infected by an index child along the essential transmission motif (solid, color) but does not include all relationships that may contribute to transmission (dashed black).

Infections in index children are defined to begin on day $t = 1$ due to either mOPV or WPV.
exposure on day \( t = 0 \). Incidence is determined by the dose response model,

\[
P_{\text{index}}(\text{infected at } t) = \begin{cases} 
p_x P(\text{infection}|\text{dose}, N_{\text{Ab, index}}) & t = 1 \text{ days} \\ 0 & t > 1 \text{ days} \end{cases}
\] (1)

where \( p_x \) (serotype \( x = 1, 2, 3, p_x \in (0, 1] \)) is a setting-specific dose response modifier that accounts for non-immunological host factors such as non-polio enterovirus infection or enteropathy that can reduce the probability of shedding [25,81,82], and the second term is defined in eq. (S5). The prevalence for \( t > 0 \) after exposure is given by:

\[
P_{\text{index}}(\text{shedding at } t) = P_{\text{index}}(\text{infected at } t = 1) P(\text{shedding at } t|N_{\text{Ab, index}}; \text{infected at } t = 1),
\]

where the first term is eq. (1) and the second is the shedding duration model in eq. (S1). Family members are infected with probabilities determined by the dose response model, the size of the fecal dose, and the amount of virus shed by the index child. Daily incidence derives from exposure to index shedding as:

\[
P_{\text{family}}(\text{infected at } t) = P_{\text{family}}(\text{transmission at } t|\text{index shedding}) 
\times P_{\text{index}}(\text{shedding at } t|N_{\text{Ab, index}}; \text{infected at } t = 1),
\]

with

\[
P_{\text{family}}(\text{transmission at } t|\text{index shedding}) = \beta(t) \prod_{t' = 1}^{t-1} (1 - \beta(t')) 
\]

\[
\beta(t) = 1 - \left(1 - P(\text{infection}|\text{dose}(t), N_{\text{Ab, family}})\right)^{D_{ij}}, 
\]

\[
dose(t) = T_{ij} \times (\text{index concentration}(t)|N_{\text{Ab, index}}),
\]

where \( P_{\text{family}}(\text{transmission at } t|\text{index shedding}) \) is the family member incidence on day \( t \) given contact with a shedding index child, \( \beta(t) \) is the infection probability determined by the dose response model, \( D_{ij} \) is the interaction rate for an index child and family member pair (average number of fecal-oral doses per day), \( T_{ij} \) is the fecal-oral dose (micrograms of stool), and index concentration (CID50 per gram) is given by the fecal concentration model in eq. (S4). Family
member prevalence follows from convolving daily incidence with the shedding duration distribution:

\[
P_{\text{family}}(\text{shedding at } t) = \sum_{t' = 1}^{t} \left( P_{\text{family}}(\text{infected at } t') \times P(\text{shedding at } (t - t')|N_{\text{Ab,family}}; \text{infected at } t') \right).
\]

The model assumes all transmission to extrafamilial contacts occurs only through family members of index children, depending on contact dose response and fecal exposure to and the amount shed by the contacted family member. Daily incidence derives from exposure to family contact shedding as:

\[
P_{\text{extrafamilial}}(\text{infected at } t) = \sum_{t' = 1}^{t} \left( P_{\text{extrafamilial}}(\text{transmission at } t|\text{family shedding since } t') \times P_{\text{family}}(\text{infected at } t') P(\text{shedding at } (t - t')|N_{\text{Ab,family}}; \text{infected at } t') \right),
\]

with

\[
P_{\text{extrafamilial}}(\text{transmission at } t|\text{family shedding since } t') = \beta(t - t') \prod_{t'' = t'}^{t-1} (1 - \beta(t'' - t'))
\]

and

\[
\beta(t - t') = 1 - (1 - P(\text{infection}|\text{dose}(t - t'), N_{\text{Ab,family}}))^{D_{fe}},
\]

\[
dose(t - t') = T_{fe} \times (\text{family contact concentration}(t - t')|N_{\text{Ab,family}}), \tag{2}
\]

where \(D_{fe}\) is the interaction rate for a family member and extrafamilial contact pair, \(T_{fe}\) is the fecal-oral dose, and \((t - t')\) is the interval since the family contact became infected. The convolution over family contact incidence accounts for all the times at which family contacts can become infected. Extrafamilial contact prevalence follows from convolving daily incidence with the shedding duration distribution:

\[
P_{\text{extrafamilial}}(\text{shedding at } t) = \sum_{t' = 1}^{t} \left( P_{\text{extrafamilial}}(\text{infected at } t') \times P(\text{shedding at } (t - t')|N_{\text{Ab,extrafamilial}}; \text{infected at } t') \right).
\]
Local reproduction number. We defined the local reproduction number ($R_{loc}$) as the expected number of extrafamilial contacts infected by an index child due to transmission along the index-family-extrafamilial essential transmission motif,

$$R_{loc} = p_{ie} N_e,$$

$$p_{ie} = \frac{\sum_{t=1}^{\infty} P_{\text{extrafamilial}}(\text{infected at } t)}{\sum_{t=1}^{\infty} P_{\text{index}}(\text{infected at } t)},$$

where $N_e$ is the average number of extrafamilial contacts and $p_{ie}$ is the total probability that an index child transmits through an older sibling to an extrafamilial contact in another household, as determined by the ratio of total incidences in eqs. (1) and (2). $R_{loc}$ describes the first generation of house-to-house transmission following infection of an index child (Fig 5B).

Within close-knit communities where all households have similar demographic, behavioral, and immunological patterns, $R_{loc}$ provides a lower bound on the total force of infection because it does not include all possible transmission routes (Fig 5C). Across large, heterogeneous communities, the a priori relationship between $R_{loc}$ and the true average force of infection across all contacts is unclear. The model can be extended to describe any set of relationships—for example, a family member may have many more socially-distant contacts that receive smaller doses less often—but the model complexity that needs to be constrained increases rapidly with the number of relationships. For these reasons, this iteration of the model cannot make predictions about the absolute probability or severity of outbreaks, for which model specification is critical [47–51]. Rather, $R_{loc}$ is a useful threshold parameter for categorizing outbreak risk with data from contact-tracing studies.

Calibration. While the data on within-host aspects of polio infection showed remarkable coherence across studies from different eras and settings, this is not the case for literature on community transmission of poliovirus. The eighteen transmission studies reviewed by Tebbens et al. [24] exhibit varying thoroughness in their reporting pre-exposure immunity and contact relationships. In lieu of a comprehensive review, we based our transmission model on specific studies capable of identifying important model parameters. The studies took place in the United States between 1953 and 1960 [35,36] and India between 2003 and 2008 [37]; all had large sample sizes, carefully reported demographic and social contact attributes, and provided sufficient information to infer pre-exposure OPV-equivalent immunity (either directly through vaccination histories or serostatus, or indirectly via shedding duration). The fraction of subjects positive for poliovirus after
OPV challenge or WPV exposure was given by the number of subjects shedding in stool or recently seroconverted divided by the number tested. Additional information about calibration methods are provided in Supplement.

We assumed that the serotype-specific dose response parameters (Table 1, eq. (S5)) are independent of setting. The setting-specific free parameters are the pre-challenge OPV-equivalent antibody titers for each subject type ($N_{\text{Ab}}$), the average fecal-oral dose (micrograms of stool ingested per interaction, $T_{fi}$ and $T_{fe}$), the interaction rates (number of fecal-oral contacts per day for each person-to-person pair, $D_{fi}$ and $D_{fe}$), the setting-specific dose response modifiers ($p_x$), and the typical number of close extrafamilial social contacts ($N_e$). The interaction rate and fecal-oral dose parameters are not separately identifiable from the available data, and so we fixed the index-to-family-member interaction rate to once per day ($D_{fi} = 1$) and assumed that fecal-oral dose is independent of relationship type ($T_{fi} = T_{fe}$).

From the Sabin transmission study conducted in Houston 1960, we calibrated the serotype-specific dose response parameters, and the fecal-oral dose and family/extrafamilial interaction rate representative of a typical endemic setting with low socioeconomic status in the pre-elimination United States. Additional study-specific parameters described OPV-equivalent immunity and trial-to-trial variation in post-vaccination shedding in index children. Briefly, children aged 2 to 18 months were enrolled to receive a dose of mOPV. Weekly stool samples were collected from the vaccine-recipient index children, their siblings (under age 15 years; average age 4 years), and primary extrafamilial social contacts of siblings. The majority of index children had prior serological immunity either due to maternal antibodies or prior IPV vaccination. Pre-challenge serology was not presented for siblings or contacts. The authors observed no significant differences in shedding by IPV immunization history or pre-challenge serologic immunity. Family members and contacts five to nine years of age shed significantly less from transmission, and there was essentially no shedding in subjects older than ten years of age (Supplement). From joint calibration across the three mOPV trial arms (Fig 3A), we inferred that children under five years of age who shed poliovirus, regardless of position in the transmission chain, had OPV-equivalent antibody titers of $N_{\text{Ab}} = 1$, and the fraction of infants shedding one week after receiving mOPV was high: type 2, 0.92 (0.85, 1.0), type 1, 0.79 (0.70, 0.88), and type 3, 0.81 (0.71, 0.91). Thus it is likely that most children under five had no experience with WPV. (See Supplement for additional details.) From the differences in transmission by serotype in this immunologically-naive population, we estimated the infectiousness of each serotype (shown above in Table 1). The estimated fecal-oral dose was microscopic at 5 (1, 31)
micrograms per day ($\mu$g/day), and the estimated interaction rate in a family member and extrafamilial contact pair—the average number of fecal-oral exposures per day—is 9.0 (2.6, 46), possibly reflecting higher rates of social interaction in peer versus infant-sibling pairs [83].

**Figure 6. Transmission model calibration.** Each study measured the amount of transmission from index children in different ways. (A) Houston 1960: fraction of subjects shedding each week after mOPV challenge and subsequent transmission. (Color by subject type; weekly data MLE and 95% CI, dot-and-whiskers; model MLE and 95% CI, lines). Eight free parameters are jointly identified across the nine calibration targets. (B) Louisiana 1953–1955: incidence in family contacts of index children naturally infected by WPV, measured by seroconversion approximately 30 days after the index child became infected. Three free parameters are jointly identified by the five calibration targets. (C) Uttar Pradesh & Bihar 2003–2008: mean prevalence of WPV measured in close contacts after the onset of paralysis in index children. One free parameter is jointly identified by the two calibration targets. See Supplement for additional information about model fit.

From the WPV transmission study conducted in Louisiana from 1953–1955 [35], we calibrated WPV dose response and the age-dependence of the fecal-oral dose, under the assumption that the fecal-oral dose between index children and older siblings was the same as in Houston 1960. Briefly, Gelfand et al enrolled families with newborn children to undergo monthly surveillance for
naturally-acquired polio infections. Whenever a newly-infected index child was identified, household contacts were tested for subsequent polio infection, most reliably through evidence of seroconversion. This measure of incidence was reported for siblings and parents, stratified by serostatus and age relative to the index child. We assumed that the OPV-equivalent antibody titer of seronegative subjects was \( N_{\text{Ab}} = 1 \), and we reconstructed from the published serological data that the median seropositive titer was \( N_{\text{Ab}} = 93 \) in this naturally-immunized population. Joint calibration of incidence thirty days after index infection across the five reported index-family relationships (Fig 6B) confirmed the expected outcome that WPV is more infectious than any Sabin strain (Table 1). We inferred that the fecal-oral dose transmitted from index children to adults was 26 (16, 41)% of that passed to siblings under age five; a similar age-related decline in fecal-oral dose was inferred with this same model for a recent Sabin 2 transmission study in Bangladesh [38]. The estimated fecal-oral dose transmitted from older index children to younger siblings was 46 (26, 104)% of the reverse.

To estimate an upper-bound for fecal-oral dose in regions of extremely high polio transmission intensity [84], we examined WPV surveillance data from 2003–2008 in India and reported by Grassly et al [37]. The authors examined the fraction of stools positive for WPV from children who were close contacts (household members or playmates) of paralytic WPV cases (mostly from Uttar Pradesh (UP) and Bihar). Contacts with low immunity (0–2 reported tOPV doses) and high immunity (6+ reported doses) were grouped for analysis. They estimated that 51% (16, 84)% of low immunity and 12 (8, 16)% of high immunity contact stool samples were positive for WPV when sampled once during the ten weeks after paralysis of the index child. For our model, we assumed that the high immunity cohort had an OPV-equivalent antibody titer of \( N_{\text{Ab}} = 512 \), corresponding to their estimate of an eleven day mean shedding duration, and that the low immunity cohort had \( N_{\text{Ab}} = 1 \) in this setting known for low tOPV efficacy [85]. Given the assumptions, we inferred from joint calibration to both targets (Fig 6C) that the fecal-oral dose transmitted from index children to contacts under five was 230 (2, 1800) \( \mu g \)/day, roughly fifty times higher than in Houston 1960.

**Additional assumptions.** The calibration studies did not report sufficient information to constrain the average number of close extrafamilial contacts \( (N_e) \), and only the Houston study provided information about the family member to extrafamilial contact interaction rate \( (D_{fe}) \). Except when explicitly exploring sensitivity to these parameters, we made the following assumptions. For Houston/Louisiana, we assumed that the typical number of extrafamilial contacts is \( N_e = 4 \) (3, 5), reflecting the average number of close friends in American childhood social
networks [86]. For UP and Bihar, we assumed \( N_e = 10 (8, 12) \), based on scaling Houston in proportion to the two-to-three times larger typical classroom sizes [87,88] and population densities [89,90] in northern India. For all settings, the value for the family member and extrafamilial contact pair interaction rate \( (D_{fe}) \) estimated from Houston was used. To simplify the presentation of results below, we chose to ignore adults because calibration showed that changes in childhood immunity from vaccination policy choices have larger effects on immunity than waning (Fig 1C and Fig 4) and that, after controlling for immunity, adult family members of infected children remained less likely to become infected than siblings (Fig 6B) [35,38].

Results

Fig 7 summarizes our within-host model for the effects of immunity on correlates of transmission and how typical immunity levels relate to specific vaccination schedules. The shedding index (Fig 7A) is the expected total amount of virus shed per gram of stool after mOPV challenge. For a typical healthy child under five years of age—averaged over vaccination timing and waning—each of the first three doses of OPV increases the OPV-equivalent antibody titer by roughly a factor of eight and decreases the expected amount of virus shed by a factor of ten. For settings with low OPV effectiveness [25,81,82,85,91], we found from calibration to data from Uttar Pradesh and Bihar [37] six or more doses were required to yeild OPV-equivalent immunity similar to that of three doses in healthy trial subjects. In our model, IPV boosting and OPV doses after the first three maintain maximum immunity. The heterotypic protection against type 2 from bOPV immunization is comparable to that of a single homotypic dose but does not accumulate with multiple doses. We inferred from the trial arms reviewed that the OPV-equivalent immunity of IPV-only is at most comparable to heterotypic immunity from bOPV, but we expect that the true impact is closer to none—the trial arms that showed the highest immunity (Fig 1) likely included some incidental IPV boosting, with larger effects in older [42,62,66,70] vs. younger [54,58,62] subjects in OPV-using countries, and negligible effects in older subjects in OPV-free countries [68,92]. Susceptibility is also strongly impacted by immunity, with the expected fraction shedding after Sabin 2 challenge dropping below half at all relevant doses for \( N_{Ab} \geq 64 \) (Fig 7B).

Our waning model (eq. (S6), Fig 4) predicts that without reinfection, fast waning causes a decline from maximum OPV-equivalent antibody titers (\( N_{Ab} = 2048 \)) to typical three-dose healthy child immunity (\( N_{Ab} = 512 \)) in 5 (4, 7) months, and an additional 4 (2, 10) years to fall to typical
two-dose immunity ($N_{Ab} = 64$). However, the model also predicts that it takes an additional 45 (15, 160) years to fall to the equivalent of one-dose childhood immunity ($N_{Ab} = 8$) and that residual immunity persists for life regardless of serostatus, as has been suspected previously [24,93]. This result disagrees with the conclusions of Abbink et al [69]. They argued from the lack of correlation between serological boosting responses and shedding duration after OPV challenge that memory immunity in seronegative elderly does not protect against poliovirus shedding, but the study lacked a control group of never-exposed subjects to contrast deeply waned and truly naive immunity. As seen through metastudy, the OPV-equivalent immunity of the Abbink et al seronegative elderly cohorts is similar to that of children who have received one dose of OPV. For heterotypic immunity against type 2 from bOPV, we predict that protection from shedding will be lost 13 (9, 22) months after bOPV vaccination is stopped [94].

Fig 8 shows maximum likelihood estimates from our transmission model for the local reproduction number of WPV, $R_{loc}$ (eq. 3), as functions of immunity and daily fecal-oral dose (Fig 8A), and fecal-oral dose and the number of extrafamilial contacts (Fig 8B). The value of $R_{loc}$, a measure of the average force of infection in a community, depends linearly on the number of extrafamilial contacts, but varies across four orders of magnitude due to strong effects of immunity and dose. Assuming one fecal-oral exposure per day (see Methods: Transmission model: Calibration), the physiological range for the average fecal-oral dose maxes out at two milligrams of stool, corresponding to the upper bound of our estimate from Uttar Pradesh and Bihar in 2003–2008. When all children have typical three-dose childhood immunity or more ($N_{Ab} \geq 512$), we estimated
$R_{\text{loc}} < 1$ over the entire physiological range, and thus that WPV persistence is impossible under universal tOPV immunization. In the absence of immunity, WPV epidemics are possible in all settings where sanitation practices permit the ingestion of roughly one microgram of stool per day or more.

Figure 8. WPV local reproduction number depends on immunity, sanitation, and contact network size. (A) Local reproduction number vs. immunity and fecal-oral dose (assuming twelve extrafamilial contacts per index child and that everyone has equal immunity). (Colormap, $R_{\text{loc}}$; dashed lines, force of infection category boundaries; HL, Houston/Louisiana; UP, Uttar Pradesh and Bihar.) (B) Local reproduction number vs. fecal-oral dose and number of extrafamilial contacts (assuming all are immunologically-naive; legend as in panel A).

We identified three categories describing the force of infection in different settings: low, where the fecal-oral route alone cannot sustain WPV transmission ($R_{\text{loc}} < 1$ for all $N_{\text{Ab}} \geq 1$); moderate, where WPV epidemics can occur in immunologically-naive communities but not where at least one-dose OPV-equivalent immunity is common ($R_{\text{loc}} \geq 1$ only for $N_{\text{Ab}} < 8$); and high, where WPV can persist despite at least one-dose OPV-equivalent immunity in everyone ($R_{\text{loc}} \geq 1$ when $N_{\text{Ab}} \geq 8$ but less than a protective threshold).

Fig 9 shows the dependence of the local reproduction number on poliovirus strain and immunity for example low, moderate, and high force of infection settings. In low force of infection settings, epidemic transmission of any strain cannot occur without contributions from the unmodeled oral-oral transmission route. This result supports the long-held hypothesis that oral-oral transmission is critical in settings with good sanitation, supported by many observations that IPV alone—an effective intervention against oral shedding [39–43]—can block transmission and prevent outbreaks from importation in communities with high socioeconomic status [8,43,95]. In moderate force of infection settings (such as Houston 1960 [36], Louisiana 1953–1955 [35], or Matlab, Bangladesh 2015 [36]), immunologically-naive populations can support WPV epidemics, but $R_{\text{loc}} \lesssim 1$ for the Sabin strains, and one-dose OPV-equivalent immunity ($N_{\text{Ab}} = 8$) is sufficient to block...
epidemic transmission of all strains. This result is consistent with the historical experience in middle-
and high-development countries that WPV elimination rapidly follows the introduction of OPV 
and that circulating vaccine-derived poliovirus (cVDPV) outbreaks are 
unknown outside of isolated communities with atypical immunological and social 
conditions.

**Figure 9. Effects of poliovirus strain on the local reproduction number.** $R_{loc}$ vs. of 
infectiousness (HID50) and OPV-equivalent antibody titer for low (fecal-oral dose $T_{f} = 0.5 \mu g/day$ 
and number of extrafamilial contacts $N_e = 3$), moderate (Houston 1960, $T_{f} = 5 \mu g/day$ and 
$N_e = 4$), and high (UP and Bihar, $T_{f} = 230 \mu g/day$ and $N_e = 10$) force of infection settings. 
(Colormap, $R_{loc}$; dashed lines, MLE for the HID50 of each strain (Table 1).)

In high force of infection settings (such as UP and Bihar 2003–2008), reinfection of previously 
immunized people can permit community-wide epidemics if typical immunity is below a threshold 
level. In the example shown, one-dose OPV-equivalent immunity ($N_{Ab} = 8$) has little or no impact 
on $R_{loc}$ for any poliovirus strain, and WPV elimination requires $N_{Ab} > 64$ for all. This result that 
WPV could persist despite $N_{Ab} > 8$ for most children in UP and Bihar 2003–2008 is supported by 
ersosurveillance. Prior to WPV elimination, the endemic dynamics of natural infection and 
vaccination conspire to maintain typical immunity levels near $R_{loc}^{(WPV)} \approx 1$, and thus the Sabin 
strains must have $R_{loc}^{(Sabin)} < 1$, with Sabin 2 highest and Sabin 3 lowest. This result is consistent 
with the historical experience that vaccine-derived outbreaks have only been observed after genetic 
reversion has restored WPV-like properties where the WPV serotype has been eliminated, and that type 2 cVDPV are most common. However, if poliovirus is re-introduced after 
elimination into a high force of infection setting with insufficient immunity, our model predicts that 
epidemic dynamics will be similar for all strains: $R_{loc}^{(Sabin)} \approx R_{loc}^{(WPV)} > 1$ is determined by the 
number of social contacts and is insensitive to differences in infectiousness of the Sabin strains.

Our results above, combined with the observation that cVDPV outbreaks have only been 
observed at rates of roughly one per year per 250 million children at risk under fifteen years of age.
indicate that settings where the force of infection for the Sabin strains is high have been rare. To evaluate how community susceptibility to Sabin 2 transmission will change due to anticipated vaccination policy changes after WPV eradication [94], we considered four scenarios for childhood immunity against type 2 poliovirus in Fig 10A. The tOPVx3 scenario describes pre-cessation populations where all index, family member, and extrafamilial contacts had achieved maximum immunity prior to waning. The bOPV & tOPVx3 scenario applies in the first two to three years after type 2 cessation, when birth spacing [104] is such that the likely index child in a family has only received bOPV (and possibly IPV) but older family members and their extrafamilial contacts have had tOPV. The bOPV scenario applies when two or more children in a typical family are born after type 2 cessation, and the naive scenario applies in settings where all OPV immunization has stopped. Prior and up to a few years after type 2 cessation, $R_{\text{loc}}^{(\text{Sabin 2})} < 1$ almost everywhere. However, our model predicts that $R_{\text{loc}}^{(\text{Sabin 2})} > 1$ will be common when typical families have more than one child born after type 2 cessation, in settings where hygenic practices are comparable to those of UP and Bihar in the years preceeding WPV elimination. Some moderate transmission settings may also become susceptible to Sabin 2 outbreaks once all OPV vaccination is stopped.

To relate local reproduction number to data that can be collected in the field, Fig 10B shows our maximum likelihood estimates for the fraction of index children, family members, and extrafamilial contacts that shed after mOPV2 challenge of the index child. In well-protected communities ($R_{\text{loc}} \ll 1$), the model predicts little to no measurable transmission from index children infected with Sabin 2, but when $R_{\text{loc}} \gg 1$, Sabin 2 transmission from index children to unvaccinated contacts will be nearly indistinguishable from WPV [24,35,37].

Fig 11 shows the sensitivity of the local reproduction number in immunologically-naive settings to extrafamilial social distance, measured in terms of the fecal-oral dose ($T_{fe}$) and the family member to extrafamilial contact interaction rate ($D_{fe}$). In moderate force of infection settings such as Houston 1960, $R_{\text{loc}}$ declines rapidly with increasing social distance even in the absence of immunity. Relative to the calibrated parameters that describe transmission among close contacts, a ten-fold reduction in either fecal-oral dose or interaction rate reduces $R_{\text{loc}}$ from near 1 to less than 0.1. In moderate force of infection settings, significant transmission requires regular, undiluted contact, and so Sabin 2 is unlikely to spread outside of the communities it is delivered to. However, in high force of infection settings such as UP and Bihar 2003–2008, $R_{\text{loc}}$ can remain above 1 across two orders of magnitude in fecal-oral dose or interaction rate—and above 0.1 across three. Under these conditions, transmission does not require undiluted fecal-oral contact, and Sabin 2 can escape
Figure 10. The effects of vaccination policy on Sabin 2 transmission for four immunity scenarios: tOPVx3 (index and family/extrafamilial child $N_{Ab} = 512$), bOPV & tOPVx3 (index $N_{Ab} = 512$ and family/extrafamilial $N_{Ab} = 256$), bOPV (index $N_{Ab} = 8$ and family/extrafamilial $N_{Ab} = 2$), and naive (index and family/extrafamilial child $N_{Ab} = 1$). (A) Local reproduction number vs. fecal-oral dose and number of extrafamilial contacts. (Colormap, $R_{loc}$; dashed lines, force of infection category boundaries from Fig 8B; symbols, example low, moderate, and high force of infection settings.) (B) Maximum likelihood estimates of the fraction shedding for each subject type after mOPV2 challenge in young index children, for each immunity scenario and example force of infection setting in panel A.
local communities via social interactions that take place only a few times per year.

Figure 11. Effects of increasing social distance on the local reproduction number of Sabin 2 in immunologically-naive populations ($N_{Ab} = 1$). Local reproduction number vs. family member to extrafamilial contact interaction rate and fecal-oral dose, for moderate ($N_e = 4$, $T_{if} = 5 \mu g/day$, $T_{fe} \leq 5 \mu g/day$) and high ($N_e = 10$, $T_{if} = 230 \mu g/day$, $T_{fe} \leq 230 \mu g/day$) force of infection settings. (Colormap, $R_{loc}$; symbols, example parameter values from Fig 10.)

Discussion

We have shown how the effects of immunity on individual-level correlates of transmission interact with sanitation and local interfamilial relationships to determine community susceptibility to poliovirus transmission. We found that the local reproduction number is a useful threshold statistic for characterizing the force of infection. The highest typical levels of OPV-equivalent immunity in our model predict $R_{loc} < 1$ for all strains in all settings. In low and moderate force of infection settings, the Sabin strains have $R_{loc}^{Sabin} < 1$ due to attenuated infectiousness relative to WPV (Fig 9), and thus significant person-to-person Sabin transmission is unlikely regardless of population immunity. However, in high force of infection settings with low population immunity—a situation than can only exist in the absence of endemic transmission and OPV vaccination—our model predicts that the Sabin strains will have unprecedented transmission potential if re-introduced, approaching that of WPV and with highest risk for Sabin 2.

Other published mathematical models known to us have explored the effects of immunity on Sabin transmission [20,26,31,32]. Despite substantial methodological differences, all are in agreement that the Sabin strains will have reproduction numbers above one in high force of infection settings with low population immunity. In addition to novel results for dose response and waning, the key innovation of our work is its direct connection from correlates of transmission that can be measured by stool surveys to assessment of community susceptibility (Fig 10). A recent application
of this model to a field transmission study in Matlab, Bangladesh [38] found that moderate force of infection conditions exist in a low-income, high-density community in the developing world where comprehensive maternal and child health care and improved sanitation systems are in place [105]. The key limitation of our model is that, while it can predict when the outbreak risk from OPV vaccination is negligible, it cannot address the absolute probability, severity, or geographic scope of outbreaks when they are possible without incorporating additional structural assumptions and calibration data about socially-distant transmission. We discuss the relevance of our results for interpreting the history of and implications for vaccination policy in the polio eradication endgame [94] below.

Before polio vaccination, most people were immunized against subsequent polio infection by natural exposure to WPV at young ages. The Sabin strains dramatically lowered the burden of paralytic disease by producing unprecedentedly high levels of immunity and displacing WPV. OPV cessation is intended to eliminate the residual disease burden caused by the Sabin strains [5, 18], but stopping OPV vaccination will reduce global immunity against poliovirus transmission to unprecedentedly low levels.

Many highly-developed countries with good sanitation and smaller family sizes have maintained polio elimination solely through the routine use of IPV [7,106,107]. Although IPV alone has little to no impact on susceptibility or shedding in stool (Fig 7), our results show that the fecal-oral route alone is incapable of supporting epidemic transmission in low force of infection settings (Fig 8). When the oral-oral route is required to permit significant transmission, IPV can prevent outbreaks by reducing oral shedding [34,39–42]. The Netherlands is an example of a country where IPV alone has been sufficient. In 1978 and 1992, there were outbreaks of WPV, but virus was found almost exclusively within high-risk groups who refused vaccination and no evidence of circulation in the well-vaccinated population was found [95,108–110]. Furthermore, many countries that could not have eliminated WPV with IPV alone a few decades ago can now be adequately protected. The United States is an example. While there is some evidence that IPV alone could reduce WPV transmission among middle- and upper-middle class families in 1960 [40], IPV vaccination of subjects with no prior exposure to live poliovirus had no impact on transmission for both Sabin and WPV strains in communities with low socioeconomic status [36,111]. However, since 2000, the United States has only used IPV [106] and yet has remained polio-free in all vaccinated populations [100] despite extensive international connections and cross-border mixing with OPV-using countries [112].

The 2013 WPV outbreak in Israel shows the limits of IPV to prevent transmission. Eight years
after Israel switched from using both OPV and IPV to using IPV only, a type 1 WPV outbreak was tracked via sewage surveillance from February 2013 until April 2014 [113,114]. Most infections were found in children born after the switch despite 93% coverage with two or more doses of IPV and waning immunity in older people [15]. A recent model estimated that the effective reproduction number of WPV among children in the Bedouin community in which transmission was most common was 1.8 [15]; the corresponding reproduction numbers for the Sabin strains, assuming our model of infectivity, are 0.4 and below. Like the United States six decades ago, Israel in 2013 is an example of a moderate force of infection country where WPV can persist despite comprehensive IPV vaccination but the Sabin strains cannot [116–118].

In the above scenarios, OPV cessation is stable. OPV can be used to interrupt outbreaks of WPV or imported (WPV-like) cVDPV, and the persistence of vaccine-derived strains is unlikely within (Fig 10A) or outside (Fig 11A) the outbreak response zone. However, in high force of infection settings with low immunity, Sabin transmission to unvaccinated contacts within outbreak response regions will be common (Fig 10) and significant transmission to socially-distant contacts can occur (Fig 11B). In these settings, OPV cessation is inherently unstable—if poliovirus is re-introduced, there is no guarantee that transmission can be stopped and new cVDPV prevented without restarting OPV vaccination in all high force of infection settings.

The conclusion that global OPV cessation is unstable follows from the fact that doses acquired via fecal-oral exposure can be much higher in the developing world than they were in the countries where Sabin OPV was first studied and where OPV cessation has already been successful (Fig 8). The time when instability will reveal itself is uncertain. Our model predicts that two or more children per family born after cessation are required to support Sabin 2 outbreaks in most high force of infection settings (Fig 10). The median birth spacing in most bOPV-using countries is 24–36 months [104]. Thus, we predict that between early 2018 and mid-2019, the risk of establishing type 2 cVDPV will increase substantially in many regions of the developing world that have not received post-cessation mOPV2 campaigns. The cross-immunity from bOPV against type 2 (with or without IPV) does not alter this conclusion.

Our estimate of 2 to 3 years to increased cVDPV2 risk upon Sabin 2 re-introduction is consistent with predictions from other models [20,26,32,33] and is compatible with the known epidemiology of cVDPV2 outbreaks. The first known example of widespread circulation of Sabin 2 after a small release took place in Belarus in 1965, but was only confirmed as such in 2003 [119]. Two years after a local experiment in type 2 OPV cessation, tOPV given to forty children likely spread...
Sabin-derived poliovirus throughout a city of 160 thousand people for at least ten months. In northern Nigeria, after widespread vaccine refusal in 2003 and 2004 [120], restoration of tOPV vaccination seeded twelve independent type 2 Sabin-derived, including the largest known outbreak of cVDPV2 in history [45].

The introduction of IPV in routine immunization globally between 2014 and 2016 aimed to provide protection against poliomyelitis to children born after OPV cessation [121]. But without substantial improvements in sanitation, IPV supply [7], and routine immunization coverage, IPV alone is insufficient to protect against poliovirus circulation in all settings. In pursuit of high vaccine efficacy with low virulence [4,122], Sabin selected strains that are 1,000–10,000 times less likely to cause paralysis than WPV [5], but only four to ten times less infectious (Table 1). Absent population immunity, the infectiousness allows transmission and genetic evolution restores virulence [44], and so the Sabin vaccines are also insufficient [20,123,124]. To secure polio eradication for all times and in all conditions, improved vaccines that produce infection-blocking immunity without the risks of Sabin OPV are required. Genetically-stabilized, engineered live vaccines are in development and promise the benefits of Sabin OPV without the risks [125–127], and adjuvanted IPV may provide a complementary route to a new effective vaccine [128].

Regardless of the challenges detailed above, Sabin OPV vaccination is always preferable to natural infection by WPV or cVDPV. Thus, mass vaccination with OPV remains the most effective intervention to eliminate poliovirus transmission [3], and the continued use of mOPV2 in regions experiencing type 2 outbreaks is warranted [18] despite concerns about poliovirus containment [19]. For risk mitigation, our model shows the value of healthy contact stool surveillance. The fraction of vaccine recipients and unvaccinated contacts shedding is a direct probe of population immunity and the local force of infection, and our results provide a rubric to categorize the risk of subsequent transmission. To go from outbreak risk categorization to risk quantification, ongoing work to better understand the relationships between local and non-local transmission are needed [129–131].

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Competing Interests

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Supplement: Assessing the stability of polio eradication after the withdrawal of oral polio vaccine

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S1 Parameter table.

The values of all parameters used in the model, both from calibration and in the Results presentation, are shown in Table S1.
| Component | Equation | Parameter | Value (Range) | Meaning |
|-----------|----------|-----------|--------------|---------|
| OPV-equivalent antibody titer | - | $N_{Ab}$ | (1, 2)$^1$ | Individual correlate of immunity |
| Probability of shedding duration | (S1) | $\mu_S$ | 30.3 (23.6, 38.6) days | Sabin median shedding duration ($N_{Ab} = 1$) |
| | | $\sigma_S$ | 1.86 (1.57, 2.27) days | Sabin scale parameter |
| | | $\mu_{WPV}$ | 43.0 (35.7, 51.7) days | WPV median shedding duration ($N_{Ab} = 1$) |
| | | $\sigma_{WPV}$ | 1.69 (1.21, 1.94) days | WPV scale parameter |
| | | $\delta$ | 1.16 (1.13, 1.21) days | Median reduction per log$_2(N_{Ab})$ |
| Peak shedding vs age | (S2) | $S_{\text{max}}$ | 6.7 (5.9, 7.5) CID50/g | Maximum stool concentration at age 7 months |
| | | $S_{\text{min}}$ | 4.3 (3.5, 5.0) CID50/g | Maximum stool concentration at older ages |
| | | $\tau$ | 12 (1, 45) months | Decay time constant of peak concentration with age |
| Peak shedding vs immunity | (S3) | $k$ | 0.056 (0.01, 0.079) | Shedding reduction with log$_2(N_{Ab})$ |
| Shedding concentration vs time | (S4) | $\eta$ | 1.65 (1.26, 2.09) | Location parameter |
| | | $\nu$ | 0.17 (0.01, 0.78) | Scale parameter |
| | | $\xi$ | 0.32 (0.08, 0.71) | Time-dependent scale |
| Dose response | (S5) | $\alpha$ | 0.44 (0.29, 0.83) | Shape parameter |
| | | $\gamma$ | 0.46 (0.42, 0.50) | Immunity-dependent shape parameter exponent |
| | | $\beta_{S1}$ | 14 (3, 59) CID50 | Sabin 1 scale parameter |
| | | $\beta_{S2}$ | 8 (2, 30) CID50 | Sabin 2 scale parameter |
| | | $\beta_{S3}$ | 18 (5, 63) CID50 | Sabin 3 scale parameter |
| | | $\beta_{WPV}$ | 2.3 (0.3, 37) CID50 | WPV scale parameter |
| Waning immunity against infection | (S6) | $\lambda$ | 0.87 (0.73, 1.02) | Immunity decay exponent |
| Houston Sabin transmission (S1–6 & 1–3) | | $N_{Ab}$ | 1 | Pre-challenge immunity (all subjects) |
| | | dose | 10$^6$ | Vaccine dose [CID50] |
| | | $p_{S1}$ | 0.79 (0.70, 0.88) | Setting-specific mOPV1 modifier |
| | | $p_{S2}$ | 0.92 (1.0, 0.85) | Setting-specific mOPV2 modifier |
| | | $p_{S3}$ | 0.81 (0.71, 0.91) | Setting-specific mOPV3 modifier |
| | | $A_i$ | 12 months | Assumed age of index (infant) |
| | | $A_s$ | 48 months | Assumed age of family member and extrafamilial contact |
| | | $T_{if}$ | 5 (1, 45) µg per day | Fecal-oral dose from infant to family member under 5 years of age |
| | | $D_{if}$ | 1 per day | Interaction rate of infant to family member pairs (assumed) |
| | | $T_{fe}$ | 5 (1, 45) µg per day | Fecal-oral dose from family member to extrafamilial contact (assumed) |
| | | $D_{fe}$ | 9 (3, 46) per day | Interaction rate of infant to family member pairs |
| Louisiana WPV (as in Houston unless shown) (S1–6 & 1–3) | | $p_{Sx}$ | 1 | Setting-specific mOPVx modifier |
| | | $N_{Ab, aero(-)}$ | 1 | Pre-exposure immunity (index case and seronegative family members) |
| | | $N_{Ab, aero(+)}$ | 93 | Pre-exposure immunity (seropositive family members) |
| | | $T_{if, young}$ | 2.3 (1.3, 5.3) µg per day | Fecal-oral dose from index child to younger family member |
| | | $T_{if, adult}$ | 1.3 (0.8, 2) µg per day | Fecal-oral dose from index child to adult family member |
| UP & Bihar WPV (as in Houston unless shown) (S1–6 & 1–3) | | $p_{Sx}$ | 1 | Setting-specific mOPVx modifier |
| | | $N_{Ab,0-2}$ | 1 | Pre-exposure immunity (tOPV 0–2 doses) |
| | | $N_{Ab,6+}$ | 512 | Pre-exposure immunity (tOPV 6+ doses) |
| | | $T_{if}$ | 230 (2, 18,000) µg per day | Fecal-oral dose from infant to family member |
| Results (unless varied in figure) (S1–6 & 1–3) | | $p_{Sx}$ | 1 | Setting-specific mOPVx modifier |
| | | $A_i$ | 12 months | Assumed age of index (infant) |
| | | $A_s$ | 48 months | Assumed age of family members and extrafamilial contacts |
| | | $D_{if}$ | 1 per day | Interaction rate of infant to family member pairs (assumed) |
| | | $D_{fe}$ | 9 (3, 46) per day | Interaction rate of infant to sibling pairs (assumed) |
S2 Within-host model

S2.1 Sources of data on shedding and dose response

Almost all relevant studies on OPV shedding, acquisition, and transmission published prior to 2012 were reviewed by Duintjer Tebbens et al [1]. Digitized data on shedding duration and concentration of poliovirus in stool were taken from the Supplementary Material of Behrend et al [2], corrected where discrepancies were noticed, and studies involving bOPV were added [3–5]. Dose response data were digitized from the cited references [6–9]. The analyses are broadly inclusive of published data, but this paper does not represent a systematic review with pre-specified exclusion criteria. Whole studies and trial arms were excluded if they reported evidence of substantial unmeasured exposure to poliovirus prior to OPV challenge [10–16] or when data across serotypes could not be disaggregated [17]. We included OPV challenge studies where subjects experienced low levels of natural exposure to WPV or OPV during the study, provided published evidence showed that most of the subjects were unaffected [7,9,18,19]. A summary of all included data describing vaccination schedules, OPV challenge formulation or WPV exposure, ages, and shedding and dose response data, and possible natural exposure is provided in Table S2 [3–9,18–31]. For a deeper discussion of data quality from reviewed studies, see Duintjer Tebbens et al [1].
Table S2. OPV challenge studies included in analysis. Ages rounded to nearest month. “Live virus exposure” indicates possible uncontrolled exposure to OPV or WPV during study. More detailed information about the included and considered but excluded studies can be found in the digitized data tables available at [github.com/famulare/howPolioVaccinationAffectsPoliovirusTransmission](https://github.com/famulare/howPolioVaccinationAffectsPoliovirusTransmission).

* IPV administered at same time as OPV; † IPV administered alone but after prior OPV.

| RI schedule | RI schedule | challenge | age at challenge | location | publication date | live virus exposure | shedding duration | shedding titer | dose response | reference |
|-------------|-------------|-----------|-----------------|----------|-----------------|-------------------|------------------|--------------|-------------|-----------|
| seronegative | mOPV1 | 5 y | Netherlands | 1959 | yes | yes | no | no | Verlinde1959[20] |
| seronegative | mOPV1 | 13 m | UK | 1961 | no | yes | yes | no | Dick1961[21] |
| seronegative | mOPV1 | 12 m | UK | 1961 | no | no | yes | yes | Dane1961[22] |
| seronegative | mOPV1 | 2 y | USA | 1961 | no | yes | no | no | Horstmann1961[22] |
| unvaccinated | mOPV1 | 6 m | USA | 1962 | no | yes | no | no | Holguin1962[23] |
| unvaccinated | mOPV1 | 16 m | UK | 1966 | yes | yes | yes | yes | Henry1966[24] |
| unvaccinated | mOPV1 | 6 m | USA | 1967 | no | yes | no | no | Benyesh-Melnick1967[25] |
| unvaccinated | mOPV1 | 2 m | UK | 1981 | no | no | no | yes | Minor1981[26] |
| unvaccinated | mOPV1 | 2 m | China | 1986 | no | yes | no | no | Dong1986[26] |
| unvaccinated | mOPV1 | 7 m | USA | 1991 | yes | yes | no | yes | Onorato1991[27] |
| unvaccinated | mOPV1 | 7 m | Romania | 1997 | yes | yes | no | no | Ion-Nedelcu1997[28] |
| unvaccinated | mOPV1 | 2 m | USA | 2005 | yes | yes | no | no | Abbink2005[29] |
| unvaccinated | mOPV1 | 2 m | Egypt | 2008 | no | yes | no | no | El-Sayed2008[30] |
| mOPV1x3 | mOPV3 | 2,4,7 m | tOPV | 10 m | no | yes | no | no | Swartz2008[31] |
| tOPVxN | tOPV | 2,3,4 m | mOPV3 | 1,5,10 y | India | yes | no | no | yes | Jafari2014[32] |
| IPVx2 | IPVx3 & tOPVx3 | 2*,3,4 m | mOPV2 | 6 m | Chile | no | yes | yes | no | O’Ryan2015[33] |
| bOPVx3 | IPVx3 | 2,3,4 m | mOPV2 | 5 or 9 m | Latin America | no | yes | yes | no | Asturias2016[34] |
S2.2 Shedding duration after OPV challenge or WPV infection

For the model calibration results in Fig. 1, we aggregated all trial arms based on poliovirus strain (Sabin 1,2,3 or WPV) and pre-challenge immunization history (formulation and number of vaccine doses). We ignored differences in vaccination schedule (i.e. tOPV at 6, 10, 14 weeks of age [5] was grouped with 7, 8, 9 months [7]) and age at challenge because our goal was to model levels of immunity that describe typical conditions among children in practice—where vaccination schedules are not rigorously adhered to and age of natural exposure is not related to age of vaccination. We also ignored the exact dose (i.e. Salk vaccine [7] vs eIPV [9]) as dose effects on shedding were insignificant at this level of aggregation. Because individual-level data was not available, we could not construct proper Kaplan-Meier estimates of the survival distribution for each immunization history. Rather, we assumed that the fraction shedding at each time point for each trial arm represented an approximate survival distribution, and the aggregated distributions shown in Fig. 1 are the sample-size-weighted averages of the fraction shedding at each time point from the original papers. Thus, in rare instances where the sample sizes are small (i.e. tOPVx2), the empirical distribution is not monotonically-decreasing as is necessary for a true survival model. Given the approximate aggregated survival distributions, we estimated approximate maximum likelihood parameters of the shedding duration model using binomial maximum likelihood (assuming independent samples). We used parametric bootstrap to estimate confidence intervals.

We assumed a log-normal survival distribution for the shedding duration given infection:

$$P(\text{shedding at } t| N_{Ab}; \text{infected at } t=0) = \frac{1}{2} \left(1 - \text{erf} \left(\frac{\ln(t) - (\ln(\mu) - \ln(\delta) \log_2(N_{Ab}))}{\sqrt{2} \ln(\sigma)}\right)\right),$$  \hspace{1cm} (S1)

where $N_{Ab}$ is the OPV-equivalent antibody titer, $\mu$ is the median duration in days for immunologically-naive individuals ($N_{Ab} = 1$), $\delta$ describes the decrease in median duration with increasing immunity, and $\sigma$ describes the shape of the distribution. The median durations and OPV-equivalent antibody titers shown in Fig. 2 were estimated under this model. Figure S1 shows the model maximum likelihood estimates (MLE) and 95% confidence intervals (CI) for the shedding duration distribution at low and high OPV-equivalent antibody-titers. An earlier version of this model was published within the supplemental software of Behrend \textit{et al} [2] but was not described in that paper, and the model was used without derivation in references [33,34].

We estimated that the WPV shedding duration in immunologically-naive children was 43.0 (35.7, 51.7) days from longitudinal surveillance studies of WPV incidence, significantly longer than our estimate for shedding duration after OPV challenge, 30.3 (23.6, 38.6) days. To confirm that this estimate is not an
artifact of differences between OPV challenge and WPV surveillance study design, we examined alternative data for the time from infection to paralysis and for shedding duration after the onset of paralysis. Casey et al measured that the mean time to paralysis from WPV infection is 17 days \cite{35} and Grassly et al showed that the mean shedding duration after paralysis from WPV infection in UP & Bihar is 31 days \cite{36}. The sum, 48 days, is consistent with our previous estimate.

S2.3 Concentration of poliovirus in stool

For each trial arm that informed our concentration model \cite{4,6,20,21,28,29}, we estimated the OPV-equivalent antibody titer from the shedding duration distributions of each trial arm as above. To model the age-dependence of the concentration of poliovirus in stool, we fit an exponential model to the peak shedding concentration:

$$\log_{10}(\text{peak CID50/g|age; } N_{Ab} = 1) = \begin{cases} S_{\text{max}} & \text{age < 6 months} \\ (S_{\text{max}} - S_{\text{min}}) \exp\left(\frac{7 - \text{age}}{\tau}\right) + S_{\text{min}} & \text{age} \geq 6 \text{ months} \end{cases} \quad (S2)$$

with maximum concentration $S_{\text{max}} = 6.7 (5.9, 7.5)$, minimum concentration $S_{\text{min}} = 4.3 (3.5, 5.0)$ CID50 per gram, and time constant $\tau = 12 (1, 45)$ months. We modeled the effect of pre-challenge immunity on concentration as:

$$\log_{10}(\text{peak CID50/g|} N_{Ab}; \text{age}) = (1 - k \log_2(N_{Ab})) \log_{10}(\text{peak CID50/g|} N_{Ab} = 1; \text{age}) \quad (S3)$$
with \( k = 0.056 (0.01, 0.079) \). The poliovirus concentration timeseries peaks shortly after acquiring infection and declines slowly thereafter. To model viral load over time, following refs. \([2, 33]\), we fit a quasi-log-normal shedding profile to the age-adjusted aggregated data for immunologically-naive individuals:

\[
\text{concentration}(t) \mid N_{Ab}; \text{age} = \max \left( 10^{2.6}, \text{peak CID50/g} \mid N_{Ab}; \text{age} \right) \left( \exp \left( \eta - \frac{\nu^2}{2} - \frac{(\log(t) - \eta)^2}{2(\nu + \xi \log(t))^2} \right) \right)
\]

with \( \eta = 1.65 (1.26, 2.09) \), \( \nu = 0.17 (0.01, 0.78) \), \( \xi = 0.32 (0.08, 0.71) \), and lower bound \( 10^{2.6} \) CID50/g to reflect the minimum reported detectable shedding.

### S2.4 Dose response to OPV challenge

For each trial arm that informed our dose response model \([4\text{–}9]\), we estimated the OPV-equivalent antibody titer from the shedding duration distributions of each trial arm as above. In order to summarize data for all doses and OPV-equivalent antibody titers, we fit a beta-Poisson dose response model for the fraction shedding after receiving an oral poliovirus dose. The beta-Poisson model is based on the assumptions that a single infectious unit (measured in CID50—the amount of poliovirus required to induce a cytopathic effect in 50% of inoculated cell or tissue culture plates) is sufficient to start an infection, that multiple infectious units contribute independently to the total probability of infection, and that the probability an infectious unit survives from initial oral exposure to the site of infection is beta-distributed \([37]\). Since the model in Behrend et al \([2]\) fitted poorly at low doses and high immunity, we explored various parameterizations of the model and found that a parsimonious description of all the OPV challenge data was provided by:

\[
P(\text{infection} \mid \text{dose}, N_{Ab}) = 1 - \left( 1 + \frac{\text{dose}}{\beta} \right)^{-\alpha (N_{Ab})^{-\gamma}},
\]

where \( \alpha \) and \( \beta \) are the standard beta-Poisson parameters, \( N_{Ab} \) the OPV-equivalent antibody titer, and \( \gamma \) captures the reduction in shedding probability with increasing immunity.

We used the fitted dose response model to estimate the OPV-equivalent antibody titer after IPV boosting on children with many prior doses of tOPV in India \([3]\). The maximum likelihood estimate of the OPV-equivalent antibody titer was \( N_{Ab} = 3700 (1700, 7700) \) and not significantly different from the maximal immunity produced by tOPVx3 prior to any waning \([5]\) (\( N_{Ab} = 2048 (430, 9600) \)).
S2.5 Waning immunity against infection

For each trial arm that informed our waning model [3, 5, 9, 20, 29], we estimated the OPV-equivalent antibody titer from the shedding duration distributions of each trial arm as above.

The time interval between last immunization and mOPV challenge was either reported or estimated as follows. For individuals from tOPVx3 vaccine trials, intervals between last immunization and mOPV challenge ranged from 1 month [5] to 6 months [9]. To assess waning of tOPV-based immunity in older children, one study in Uttar Pradesh compared mOPV vaccine take rates in children 1, 5, or 10 years of age [3] who had previously received an unknown but high number of tOPV doses. To estimate the likely interval between last immunization and challenge, we assumed that children are offered up to 5 doses in the first year of life (3 RI plus 5 campaigns at 60% coverage), corresponding to roughly 2.5 months on average between last vaccination and mOPV challenge at 1 year of age. We assumed campaigns delivered 3 doses per year in ages two through four, corresponding to roughly 4 months between last vaccination and challenge at 5 years of age, and no doses after 5 years of age, corresponding to 5 years since last vaccination and challenge at 10 years of age. For this study, OPV-equivalent immunity was inferred via vaccine take rates using equation (5). Data on adult shedding after natural immunity were taken from studies in the Netherlands. From the study by Verlinde et al [20] in 1959, the average seropositive subject in the study was 20 years of age, and we assumed that their last infection was 5 years earlier at 15 years of age when maximum seropositivity was first achieved in the population. From the study by Abbink et al [29] from 2005 that measured shedding in elderly individuals upon mOPV challenge, we assumed last exposure was 45 years earlier in 1960, at roughly the year in which widespread endemic transmission ceased in the Netherlands. We included data for both seropositive and seronegative adults from the Abbink et al study because seronegative adults showed evidence of memory immunity and reduced shedding durations in comparison to immunologically-naive children.

We fit a power law waning model [38] to the OPV-equivalent antibody titers,

$$N_{Ab}(t) = \max(1, N_{Ab,1}t^{-\lambda}),$$  \hspace{1cm} (S6)

where $t$ is measured in months between last immunization and oral challenge, $N_{Ab,1}$ is the baseline immunity one month post-immunization, and the exponent is $\lambda = 0.87 \ (0.73, 1.02)$. 

\hspace{1cm} 8/17
S3  Transmission model

S3.1  Model fit

Figure S2 is an extension of Figure 6 that shows maximum likelihood estimates of the fraction shedding (prevalence) and cumulative incidence in our model for the three calibration targets. For parameters, see Table S1.
Figure S2. Model of fraction shedding and incidence for each calibration target. Maximum likelihood estimates (solid lines) are shown for each subject type (color) after mOPV in Houston or WPV in Louisiana and UP & Bihar. Dots-and-whiskers show calibration targets, and model 95% CI are shown for comparison. For Houston, we compared fraction shedding in stool to model prevalence. For Louisiana, cumulative incidence one month after index child infection. For UP & Bihar, we compared the mean fraction of close (direct personal) contacts shedding after index child paralysis (model prevalence and calibration target (mean over time) shown).
S3.2  Houston 1960

No breakdown by age was presented by Benyesh-Melnick et al. [25] for the extrafamilial contacts of the siblings. However, because the contacts are demographically similar to the siblings and age is a significant factor for poliovirus acquisition via transmission in this setting, we used age-adjusted shedding rates in this paper. To estimate the shedding fraction in the age under 5 contact cohort, we adjusted the total reported shedding counts for each serotype as follows:

\[
\text{estimated contacts shedding under 5) = (total contacts shedding) \times (fraction siblings shedding under 5)}
\]

\[
\text{(estimated contacts under 5) = (total contacts) \times (fraction siblings under 5).}
\]

The estimated counts were rounded to the nearest integer and confidence intervals presented are based on the rounded estimated counts.

Figure S3 shows more information about the age-dependence of shedding after mOPV challenge. Older index children shed slightly less after mOPV challenge than younger children for types 2 and 3 (type 1 \( p = 0.105 \); type 2 \( p = 0.016 \); type 3 \( p = 0.025 \); two-tailed Fisher’s exact test). This observation was not explored in the original paper, and we propose two possible explanations. As described in Table 1 of Benyesh-Melnick et al. [25], older index children were more likely to have received at least one dose of IPV. However, it should be noted that the original authors reported that they found no significant differences between IPV and unvaccinated index subjects, as is compatible with our metastudy. A second possibility is that stool concentrations of poliovirus are higher in young infants, and so stool culture may be more sensitive to shedding in younger children (eq. (S2)). In the family member cohorts, there were no statistically significant differences in shedding among the age groups under 12 months, 12 to 23 months, 24 to 35 months, or 36 to 59 months for any serotype. However, there was significantly less shedding in the 60 to 107 month age group relative to the 36 to 59 age group (\( p < 0.001 \) for all serotypes). As stated in the main text, shedding in siblings age 60 to 107 months (5 to 9 years) is significantly below that of ages less than 5 years for all serotypes (type 1 \( p < 0.001 \); type 2 \( p < 0.001 \); type 3 \( p = 0.002 \)). Shedding rates were very low in parents and children age 10 years and older (<2%) [25], and so it is likely the transmission was direct from index child to sibling and was not mediated by infected caretakers. Shedding due to transmission-acquired type 2 was significantly more common than for types 1 and 3, and shedding due to transmission was similar for types 1 and 3 (mean prevalence: type 1 vs type 2 \( p = 0.002 \); type 1 vs type 3 \( p = 0.33 \)). Primary extrafamilial contacts of siblings exhibited a similar pattern of increased type 2 shedding and comparable type 1 and 3.
Figure S3. Fraction shedding by cohort and age range as originally reported. Observed fraction shedding and estimated 95% binomial confidence interval for each serotype, subject type, and reported age cohort.

sheding (type 1 vs type 2 \( p < 0.001 \); type 1 vs type 3 \( p = 0.73 \)). Although the authors did not describe the relationships between siblings and extrafamilial contacts in detail, it is likely that the contacts were close friends of the siblings and were directly infected by the siblings, as the authors also describe a smaller set of more socially-distant “secondary extrafamilial contacts” who “were drawn from the neighborhoods or schools attended by the siblings” and who were infected at lower rates than the primary contacts [25].

Little information about shedding in secondary contacts was provided, except to note that, summed across all trial arms, 15 of 280 secondary contacts were positive for Sabin 2 and the highest incidence rate was 13% in the secondary extra-familial contacts of tOPV recipients. Assuming the number of secondary contacts is proportional to the number of primary contacts for each trial arm, \( n = 10 \) of the type 2 positives were in secondary contacts of tOPV recipients (13% of trial arm total) and \( n = 5 \) were in secondary contacts of mOPV2 recipients (8.5% of trial arm total).

S3.3 Louisiana 1953–1955

We calibrated model incidence to the seroconversion data reported in Gelfand et al [39]. Stool collection data was also available, but it reported lower levels of incidence. This was likely due to missing infections: the
average interval between samples was 27 days while the average shedding duration in seropositive subjects with median $N_{Ab} = 93$ is only 16 days under our model. We assumed 100% incidence of immunologically-naive index children after WPV exposure, based on the study design that reported family member incidence conditional on detection of the child’s first natural infection with poliovirus.

### S3.4 Uttar Pradesh and Bihar 2003–2008.

We calibrated our model to the estimates of mean stool prevalence after the onset of paralysis in close contacts of WPV cases reported by Grassly et al [36]. We assumed that the contact data best corresponded to family members in our model. Quote:

*During identification of healthy contacts, an effort was made to include those children with the closest contact to the individual with suspected poliomyelitis, such as siblings, playmates, or residents of the same household.* [36]

To shift our model from prevalence after infection to prevalence after paralysis, we convolved our prevalence timeseries with the time-to-paralysis distribution:

$$P_{\text{family}}(\text{shedding at } t \mid \text{index paralysis at } t = 0) = \int_0^t dt' P_{\text{family}}(\text{shedding at } t') p_{\text{paralysis}}(t - t'),$$  \hspace{1cm} (S7)

where $p_{\text{paralysis}}(t)$ was given by the histogram in Figure 2 of Casey et al [35]; the mean time from infection to the onset of paralysis was 17 days. The model was calibrated against the mean of eq. (S7) over the first 90 days after index child paralysis.

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