Review
Leveraging beneficial off-target effects of live-attenuated rotavirus vaccines

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Abstract: Following the introduction of live-attenuated rotavirus vaccines in many countries, a notable reduction in deaths and hospitalizations associated with diarrhoea in children <5 years of age has been reported. There is growing evidence to suggest that live-attenuated vaccines also provide protection against other infections beyond the vaccine-targeted pathogens. These so called off-target effects of vaccination have been associated with the tuberculosis vaccine Bacille Calmette Guérin (BCG), measles, oral polio and recently salmonella vaccines, and are thought to be mediated by modified innate and possibly adaptive immunity. Indeed, rotavirus vaccines have been reported to provide greater than expected reductions in acute gastroenteritis caused by other enteropathogens, that have mostly been attributed to herd protection and prior underestimation of rotavirus disease. Whether rotavirus vaccines also alter the immune system to reduce non targeted gastrointestinal infections has not been studied directly. Here we review the current understanding of the mechanisms underlying off-target effects of vaccines and propose a mechanism by which the live-attenuated neonatal rotavirus vaccine, RV3-BB, could promote protection beyond the targeted pathogen. Finally, we consider how vaccine developers may leverage these properties to improve health outcomes in children,
particularly those in low-income countries where disease burden and mortality is disproportionately high relative to developed countries.

**Keywords:** Rotavirus; off-target effects; neonatal; live attenuated; RV3-BB; epigenetic modulation

1. Introduction

Diarrhoea, pneumonia and malaria remain the leading causes of death among children under the age of five years [1]. Approximately five million children fail to reach the age of five every year due to treatable and vaccine preventable diseases [1]. Although efforts have been accelerated to reduce the rate of under-five mortality worldwide, an imbalance still exists between low and high income countries (HIC) [1]. In Sub-Saharan Africa, infant mortality remains high, and one in every 13 children fail to reach their fifth birthday compared to their peers in HIC [1].

Notably, diarrhoeal disease associated with rotavirus infection accounts for an estimated 128,500 under-five deaths annually, despite increasing rollout of rotavirus vaccines [2,3]. Rotavirus vaccines are key interventions to counteract the devastating effects of rotavirus associated diarrhoea [4]. To date, over 100 countries have incorporated rotavirus vaccines into their national immunization programmes [5,6]. Following implementation of rotavirus vaccination, reduction in diarrhoea-associated deaths has been observed in many countries [7]. Observational studies conducted in both high income and middle income settings suggest that rotavirus vaccines may also provide protection against severe acute gastroenteritis not associated with rotavirus [8–10]. Further understanding of the mechanisms through which new and existing vaccines induce broad-based immunity against a range of pathogens would be of considerable value to high disease burdened regions.

Both epidemiological and immunological studies suggest that certain vaccines exhibit immuno-modulatory functions that induce beneficial off-target effects against unrelated pathogens. Whilst limited data are available to explain this phenomenon, non-specific effects are in part thought to originate from epigenetic reprogramming of innate cells that subsequently lead to altered innate memory. Strong evidence on non-specific effects has been reported from live-attenuated vaccines, notably Bacille Calmette Guérin (BCG) and measles vaccines [11–14]. It is uncertain whether similar effects are applicable to rotavirus vaccines. We postulate that live-attenuated rotavirus vaccines administered orally at birth might induce beneficial non-specific effects, and confer protection to unrelated pathogens, and have therefore reviewed the available literature in this area.

2. Off-target effects of vaccines

Contrary to the traditional view that vaccines induce specific immunity to a specific pathogen, evidence has emerged over the past decades that suggests vaccines can also induce protection against non-targeted pathogens. This phenomenon has been termed ‘non-specific effects’, ‘heterologous effects’ or ‘off-target effects’ of vaccination, and from herein this article will use the latter term. Both epidemiological and immunological studies have demonstrated protective off-target effects from live vaccines whereby they reduce susceptibility to non-targeted pathogens. By contrast, non-live vaccines such as whole cell Diphtheria-Tetanus-Pertusis (DTP) may have been associated with negative effects such as slightly increased morbidity and mortality from non-targeted pathogens particularly observed in low
income-high mortality settings [12,15,16]. The World Health Organization (WHO) initially expressed reservations regarding the existence of vaccine off-target effects, largely due to data originating from observational non-randomised studies that were open to potential bias [17]. However, a systematic review conducted by WHO confirmed that measles and BCG vaccination in infancy are associated with a significantly reduced all-cause mortality from non-targeted pathogens [13], and several randomised studies have consolidated evidence that support vaccine induced off-target effects [13,15,17,18]. To date, some evidence for off-target effects have emerged from randomised studies of BCG, measles and oral polio Vaccine (OPV), with more recent evidence emerging for the live-attenuated salmonella Ty21a vaccine [46,47,50]. High-income countries such as UK and USA and low-income countries including Guinea-Bissau, The Gambia, Senegal and Sudan have all reported off-target effects of vaccination from both randomised and non-randomised studies[12,20–22].

3. Mechanisms of off-target effects of vaccination: Major lessons from Bacille Calmette Guérin

Despite the concept of off-target effects of vaccines attracting attention from scientists worldwide, the precise immunological mechanisms associated with these effects remain unclear. Nevertheless, several vaccine effect studies have investigated both innate and adaptive immune components as potential mediators of off-target effects. BCG is a widely used anti-tuberculosis vaccine and a well-studied model of vaccine induced off-target effects. BCG-induced off-target effects have been widely investigated using murine models by comparing immune responses following heterologous antigen challenge between BCG naïve and BCG vaccinated mice. Earlier studies by Spencer et al (1976) reported increased capacity of BCG stimulated macrophages in mice in reducing viral titres of influenza A virus compared with controls. A study by Starr et al further demonstrated that a single dose of BCG vaccine administered intradermally or intraperitoneally protected newborn mice from herpes simplex virus type 2 in a controlled murine model [23]. It has been suggested that shared cross-reactive epitopes between BCG and unrelated pathogens might explain the observed non-specific protective responses upon secondary exposure to the unrelated pathogens [11,24,25]. However, solid evidence is required to directly link BCG signatures to the multitude of unrelated pathogens for which off-target effects have been observed [11].

A further hypothesis is that BCG induces off-target protection via the production of cytokines that alter the inflammatory environment, thereby enabling activation of pre-existing effector or memory T cells, B cells or natural killer cells in what is called bystander effects [11]. It has been further suggested that BCG induces off-target protection via innate cell mediated production of cytokines that accelerate T follicular helper (T FH ) cell development and B cell maturation, thus enhancing a broad range of antibody responses against an array of pathogens [26–29]. Since the infant immune system is not fully developed and sufficiently experienced to generate broad pre-existing T and B cell memory, these bystander effects seem unlikely to play a major role in BCG induced off-target effects in this age group [11,30].

Additionally, Toll-like receptor 8 (TLR8) engagement with BCG is also implicated in potentiating off-target effects. BCG engagement with TLR8, which are preferentially expressed on dendritic cells, monocytes and macrophages are known to induce high levels of cytokines including interleukin 12 (IL-12) and tumour necrosis factor-alpha (TNF-α). Induction of off-target effects using this pathway appears plausible as IL-12 signalling is thought to drive differentiation and function of T FH cells that are crucial for providing help to B cells, subsequently augmenting antibody mediated immune responses. Brook et al. (2020) proposed emergency granulopoiesis as another possible mechanism linked to BCG protection against unrelated pathogens. Using a neonatal mouse model, the study demonstrated that BCG vaccination led to emergency granulopoiesis via upregulation of granulocyte colony stimulating factor (G-CSF), and that emergency granulopoiesis was associated with rapid neutrophil expansion. These findings imply that BCG-
induced granulopoiesis could potentially confer protection against microbial-induced sepsis. Similar observations were made in human neonatal studies, although the effect observed was rapid and short-lived (24hrs) [31]. If extensively studied, granulopoiesis could lessen sepsis related deaths in resource constrained settings where more than half of child mortality occurs within the neonatal period [31,32].

The concept of trained immunity is one of the most plausible mechanisms of BCG vaccine-induced off-target effects. It is based on the understanding that innate immune cells can undergo epigenetic changes following exposure to certain vaccines or microbial components [11,15]. This results in long term changes in innate immunity i.e. innate immune memory, which is at odds with prevailing thinking that has historically restricted immunological memory to the adaptive components of the immune system such as B and T lymphocytes [33–38]. Indeed, there is evidence that innate immune memory is an inherent trait in plants, animals and invertebrates and is used as a mechanism of protection [39,40]. An in vitro experimental model where primary monocytes are stimulated with potential heterologous inducers such as beta-glucan or BCG and then re-stimulated with agents such as lipopolysaccharide that act as a proxy of heterologous pathogens has been used to demonstrate innate immune training. Such studies have shown that innate immune cells can undergo metabolic, functional and epigenetic modulation leading to the acquisition of long lasting memory responses to subsequent unrelated stimuli [38].

Advancements in genome wide studies, proteomics, transcriptomics, metabolomics and epigenetics have allowed discovery of signalling pathways and signatures that may be involved in innate cell training [11,41]. Molecular pathways such as Protein Kinase B /mechanistic target of rapamycin (Akt/mTOR), Phosphoinositide/Protein Kinase B (PI3K/Akt), Nucleotide-binding oligomerization domain 2 (NOD2)-pattern receptor signalling, histone modification and NF-kB transcription factors have been highlighted as key inducers of trained immunity in monocytes. Expression of pattern recognition receptors (PRR) including TLR2, TLR3 and TLR4 as well as costimulatory molecules such as CD40, CD80 are upregulated during innate cell training. It is therefore not surprising that many studies have reported upregulation of inflammatory cytokine genes, such as IL-1β, TNF-α, IL-6, which are involved in a wide range of heterologous responses [14,42–45].

4. Off-target effects from live and non-live vaccines

Off-target effects induced by vaccines are generally characterised as beneficial or deleterious depending on the nature of a particular vaccine. Beneficial off-target effects are commonly associated with live vaccines and are thought to confer protective immunity. In contrast, deleterious effects have been reported from inactivated vaccines that are linked with increased all-cause mortality [16,19].

Off-target effects were first reported in a large cohort study conducted in Guinea-Bissau two decades ago which demonstrated increased mortality in children that received whole cell diphtheria-tetanus-pertussis (DTP) vaccine [21,22,46–50]. These effects are thought to last until the next vaccine is administered and would become beneficial if the next vaccine was a live vaccine [16]. However, the concept of deploying sequential vaccine administration to counteract negative off-target effects remains speculative considering that infants in high endemic settings are likely to be exposed to environmental or maternal pathogens, including organisms such as norovirus, enteric adenoviruses and campylobacter before the next vaccine is administered [51–53]. Whether or not these enteric pathogens would similarly enhance the developing infant immune system remains unestablished, but a similar immune-modulatory effect might not be surprising considering that these are also live organisms. Studies that could mechanistically investigate why certain vaccines appear to induce off-target effects irrespective of prevailing pathogen exposure or indeed the possibility of vaccine-pathogen co-immunomodulatory impact in high endemic countries would, to some extent, provide a better understanding on the plausibility of off-target effects in these settings. The WHO recommends concomitant
administration of rotavirus vaccine with DTP [54,55]. The impact of this co-administration on off-target effects of each vaccine requires attention in settings that are implementing these guidelines.

BCG induced off-target effects were implicated in the reduction in all-cause mortality from six controlled studies conducted in USA in the 1940-1950s [12,15,50]. Similar observations were made in a randomised controlled trial in low birth weight infants in Guinea-Bissau that reported a lower mortality rate ratio following administration of the BCG vaccine as compared to unvaccinated controls [12]. Combined findings from two randomised trials in Guinea-Bissau reported a reduced neonatal mortality ratio of 0.52 (0.33-0.82) following a birth dose of BCG as compared to no vaccine. These findings have profound implications as more than half of infant mortality in resource limited settings occurs during the neonatal period [15,16,50].

Apart from inducing specific immunity against targeted pathogens, non-live vaccines are thought to induce deleterious effects that lead to increased all-cause mortality. Some studies in high mortality settings in Africa and Asia on infants vaccinated with DTP have been linked with an increase in all-cause childhood mortality, particularly in girls [56]. A 4.33-fold increase in mortality was observed compared to controls among low birthweight neonates in Guinea randomly assigned to receive BCG vaccine at birth followed by DTP vaccine two months later. Since mortality rate is thought to be determined by the last vaccine administered, the increased mortality observed in this study was attributable to DTP vaccination [15,16]. Other vaccines associated with negative off-target effects include single hepatitis B vaccine, Pentavalent vaccine (DTP, hepatitis B virus, Haemophilus influenza type b), inactivated polio vaccine, H1N1 influenza vaccine and the malaria vaccine RTS,S/AS01 [13,56–62]. In the context of the malaria vaccine, it is clear that there is no randomised controlled Trial data hence interpretation is based on observations. Clearly, non-live vaccines are important in inducing specific immunity to their targeted pathogens, but the reported negative effects from some studies need in-depth investigation. A systematic review sanctioned by WHO on off-target effects concluded that the existing epidemiological evidence was not robust enough to support or reject deleterious heterologous effects conferred following DTP vaccination [63]. A Gambian study showed that vaccinating infants with DTP decreases both innate-pro-inflammatory and overall T cell immunity in females, providing a mechanistic explanation as to why this vaccine might increase susceptibility to other pathogens [64]. The study further speculated that downregulation of interferon gamma genes, DTP induced TLR4 inhibition as possible causes for these effects [64].

Vaccine timing is another important factor that requires consideration in order to optimise off-target vaccine effects. Indeed, it has been proposed that neonatal targeted vaccination within the first days of life as opposed to infant vaccination has the potential to modulate the infant immune system and boost immunity from birth. BCG, OPV and Hepatitis B vaccine administration during the first days of life in some countries has previously shown to be a successful and a practical means of boosting immune responses in newborns [65]. Apart from modulating the infant immune system, potential additional benefits include early protection against infections by shortening the vulnerability period where newborns are more susceptible to a range of pathogens [65]. Additionally, administration of vaccines within the first days of life appears advantageous in achieving good vaccine coverage given that healthcare services are conveniently more accessible at birth and easily implementable [65]. This would fit better in resource constrained settings where access to healthcare systems is compromised.

5. Sex differential off-target effects

The off-target effects of vaccines have been shown to be sex-differential in multiple studies. Generally, negative non-specific effects induced by non-live vaccines such as DTP appear to be more pronounced in females than males.
A study by Nohoh et al (2016) demonstrated that DTP vaccinated females had overall suppressed T-cell reactivity and innate pro-inflammatory responses and showed decreased oCD3/oCD28 stimulated type 1 cytokines responses. Contrary, live-attenuated vaccines such as smallpox, BCG and measles vaccines have shown to provide more beneficial effects in females than in males under the age of five [15,16,19,66–68]. Indeed, the same is true for the targeted effects of vaccines with females were found to elicit more robust antibody specific responses following vaccination as compared to males for multiple vaccines [66, 69]. Immunological sex differences have so far been tied to inherent variations such as hormonal and X-linked alterations that have been shown to be evident during puberty and infancy [66,68]. The mechanisms driving the observed sex differences in off-target effects of vaccination are yet to be fully elucidated. However, X-linked immune response genes and sex hormones have been suggested to play a role in these vaccine induced sex differential effects [69]. Low level expression of the X chromosome encoded TLR7 gene on B cells in influenza vaccinated males compared to females appears to contribute to the sex differential vaccine induced responses in mice [19,70]. The steroid hormone milieu has also been shown to affect vaccine induced antibody responses in males and females of both humans and mice [70]. Harnessing this information could inform development of vaccines that improve vaccine-specific responses and off-target beneficial effects in a sex-specific manner.

6. Current rotavirus vaccines

Early observations that children with secondary exposure to rotavirus were protected from moderate-to-severe rotavirus associated gastroenteritis paved the way for rotavirus vaccine development [71]. The first ever rotavirus vaccine to be approved was the human-rhesus reassortant live-attenuated tetravalent rotavirus vaccine (RotaShield®, Wyeth-Lederle, USA). RotaShield® usage was discontinued nine months after rollout in the USA in 1999 due to a reported increased risk of intussusception [5,72,73]. Currently, four live-attenuated rotavirus vaccines are WHO prequalified for global use (Table 1) and 114 countries have included rotavirus vaccines in their national or sub-national routine immunization programme. Rotarix® (GlaxoSmithKline, Belgium), a monovalent G1[P8] live attenuated vaccine and RotaTeq® (Merck, USA), a live attenuated pentavalent (G1, G2, G3, G4, [P8]) bovine-human reassortant vaccine, are currently the most widely used rotavirus vaccines [5,6]. Rotavac®, G9[P11]- (Bharat Biotec, India) and Rotasiil®, G1, G2, G3, G4, G9 (Serum Institute, India) have been introduced into the National Immunization program in India with introductions planned in Benin (Rotavac®) and Democratic Republic of Congo and Burkino Faso (Rotasiil®). The Lanzhou Lamb (G10P[12]) -and Rotavin-M1® (G1P[8])-are available on the market for use in China and Vietnam, respectively. Over 80 countries have incorporated Rotarix® into their immunization programmes, 16 countries have introduced RotaTeq® and 10 countries have introduced both Rotarix® and RotaTeq® into their routine immunisation schedules. Whilst both vaccines are administered to infants from 6 weeks of age, Rotarix® is given in two doses and RotaTeq® is administered in a 3 dose schedule [5], (Table 1).

Rotarix® is currently the most widely used rotavirus vaccine globally [6]. It was initially introduced in Mexico in 2006 and by the end of 2010, mortality due to diarrhoea had decreased by over 40% in Mexican children under the age of five years [6,74]. The decline in diarrhoea mortality could not be explained by specific vaccine induced immunity alone. Indirect protection due to a reduction in transmission of rotavirus from vaccinated children was cited as an attributable factor for the unexpectedly large decrease in diarrhoea mortality. In contrast, a study in Finland reported a lack of evidence to support reduced non-rotavirus acute gastroenteritis following vaccination with RotaTeq® [75]. The study suggested differences in geographical settings as a possible contributing factor to the conflicting results [75]. These findings support further investigations into the mechanistic explanation of off-target effects in the reported sites and the ideal population that would benefit from these effects. Prospective birth cohort studies from Malawi on mortality impact and vaccine effectiveness of sequential 13-valent pneumococcal conjugate (PCV13) and monovalent rotavirus
vaccine RV1 provided population-level evidence of infant mortality reduction further highlighting the importance of increased vaccine coverage in high-mortality settings [76].

Table 1: Current rotavirus vaccines

| Product Name | Manufacturer | Composition | Doses | Route of Administration | Regulatory status |
|--------------|--------------|-------------|-------|-------------------------|------------------|
| Rotarix®     | GSK          | G1P[8]-Live attenuated | 2     | Oral                    | WHO prequalified |
| Rotavac®     | Bharat Biologicals | G9[P11]-Live attenuated | 3     | Oral                    | WHO prequalified |
| RotaTeq®     | Merck        | G1,G2,G3,G4,P[8] -Live attenuated | 3     | Oral                    | WHO prequalified |
| Rotasil®     | Serum Institute of India | G1,G2,G3,G4,G9-Live attenuated | 3     | Oral                    | WHO prequalified |
| Rotavin-M1®  | POLYVAC      | G1P[8]-Live attenuated | 3     | Oral                    | Nationally licensed |
| Lanzhou Lamb® | Lanzhou Institute of Biological Products | G10P[12]-Live attenuated | 1     | Oral                    | Nationally licensed |

7. The need to improve the performance of rotavirus vaccines

Despite significant strides established so far following rotavirus vaccine introduction in many countries, rotavirus vaccines have generally underperformed in high mortality settings. Bergman et al. (2021) reported that rotarix prevented 93% of severe rotavirus diarrhoea cases and 52% of severe all-cause diarrhoea cases in low-mortality settings. In high mortality settings, rotarix prevented 58% of severe rotavirus diarrhoea cases and 27% of severe all-cause diarrhoea cases [77]. Underperformance of rotavirus vaccines particularly in low-income countries is of great concern that warrants development of alternative vaccines and new approaches tailored to maximise vaccine protective effects. High levels of breast milk antibodies, transplacentally transferred maternal antibodies, and a high prevalence of malnutrition and enteropathy are some of the factors that are thought to be associated with reduced rotavirus vaccine effectiveness in low income settings [78,79]. The neonatal live attenuated rotavirus vaccine RV3-BB (Biofarma, Indonesia), nonreplicating P2-VP8-P[8] and P2-VP8-P[4]P[6]P[8] subunit rotavirus vaccine (NRRV, PATH), virus-like particle vaccines (Baylor College of Medicine), inactivated whole virus vaccine (Curevac) and RNA based vaccines (CDC) [5,6,80] are alternative rotavirus vaccines currently in development (Figure 1).

The persistence of a high rotavirus disease burden in early life requires development of better vaccines that can be administered as early as the first week of life. The current licenced rotavirus vaccines are administered from 6 weeks of age, therefore missing the opportune window when the neonatal immune system is more susceptible to infections [6]. RV3-BB is an oral live naturally attenuated rotavirus vaccine developed at the Murdoch Children’s Research Institute, Australia [6]. RV3-BB is based on the human neonatal rotavirus strain RV3 (G3P[6]) associated with asymptomatic infection of the neonatal gut. Its ability to offer early protection in the presence of possible inhibitory
effects from maternal antibodies and breast milk is intriguing [81–83]. Furthermore, the immaturity of the neonatal gut microbiome and the limited intake of breast milk may present an advantage for a birth dose of rotavirus vaccine. Rotavirus VP4 [P] binding is genotype-dependent and differential expression of histo-blood group antigens (HGBA) could explain the disparity in rotavirus vaccine efficacy between populations. For P[8] infections, including with the P[8] based Rotarix vaccine, secretor-positive individuals are more susceptible to infection and have higher vaccine shedding, than observed in secretor-negative individuals. In contrast, there was no difference in vaccine take after administration of RV3-BB vaccine, based on a P[6] strain, observed by secretor status, by Lewis status or by combined secretor or Lewis status. This intrinsic feature may mean that RV3-BB has the potential to provide improved protection in Africa, where the Lewis-negative phenotype is more prevalent (20–35% in some countries) and where P[6] rotavirus strains causing disease are endemic [84,85]. [85,86].

Rotavirus vaccines are routinely administered with the first dose at 6 weeks of age. Developed from a human neonatal rotavirus strain, RV3-BB vaccine is an ideal candidate for a neonatal administration strategy, with the first dose administered at or soon after birth. RV3-BB phase II clinical trials conducted in both a high-income, low-child mortality country (New Zealand) and a low-middle income, high-child mortality country (Indonesia and Malawi) have shown that RV3-BB is immunogenic and well tolerated when administered as a three dose neonatal or a routine infant schedule [82,83,87,88]. Vaccine efficacy against severe rotavirus gastroenteritis among Indonesian infants after three doses of RV-BB in the neonatal schedule was 94% at 12 months of age and was 75% at 18 months of age [83]. In Malawian infants, anti-rotavirus IgA seroconversion was observed in 41.3% to 58.8% of participants 4 weeks post RV3-BB vaccination with three doses in the neonatal schedule. A further 62.3% to 71.9% seroconversion was noted in participants 18 weeks post RV3-BB vaccination with three doses in the neonatal arm[88].

8. Future considerations

Clearly, the concept of off-target effects of vaccines continues to evolve and important lessons are yet to be unmasked. However, there are important questions that remain to be addressed in order to utilise this intrinsic feature of vaccines to inform vaccine design. To date, limited studies have analysed for putative off-target beneficial effects of live-attenuated rotavirus vaccines either from an epidemiological or mechanistic point of view [89–91] Moreover, it is not clear whether live attenuated rotavirus vaccines incorporated into immunization programmes would influence the
performance of other vaccines known to induce off-target effects, such as BCG. Robust controlled studies that investigate how a live attenuated rotavirus vaccine administered at birth could impact off-target effects and BCG are required.

While the current understanding of mechanisms of vaccine-induced off-target effects is largely drawn from BCG studies, it would be intriguing to investigate the possible genes and signalling pathways that may be involved in influencing innate and non-targeted adaptive responses following rotavirus vaccination. Animal model studies by Lee et al (2018) demonstrated that a cold adapted live-attenuated influenza vaccine (CAIV) provided immediate protection against respiratory syncytial virus (RSV) infection. They further reported that TL3/TL7 signalling pathways were essential for these observed rapid beneficial off-target effects [92–95]. Rotavirus, like other mucosal viruses such as RSV and influenza viruses share common features despite them belonging to different virus families. For instance, infection with these viruses triggers an early innate immune response that attempts to clear infection within the mucosal sites. These viruses also commonly induce innate immunity via PRRs such as TL3, TL7 and retinoic acid-inducible gene I (RIG-I). Rotavirus similarly triggers innate sensors including RIG-I, TLR2, TL3, TLR5, TL7 and innate immune cells such as dendritic cells and natural killer cells dominantly in humans and to a lesser extent mouse models [96–99]. It is possible that some or all of the established sensors and innate cells would be plausible mediators of immunomodulatory effects by live attenuated rotavirus vaccines. Given the availability of advanced systems biology techniques at our disposal, off-target effects of live-attenuated vaccines can be explicitly studied using approaches such as epigenetics, metabolomics, proteomics and transcriptomics [11]. In vitro stimulation assays and immunophenotyping are also useful approaches to studying vaccine off-target effects [18,100].

Efforts to improve rotavirus vaccine effectiveness in low-income countries include development of next generation oral rotavirus vaccines, exploration of vaccine adjuvanticity, alternative vaccine administration schedules and alternative routes of vaccine administration such as injectable subunit or inactivated rotavirus vaccines. It is hoped that these alternative options will address some of the challenges associated with oral vaccination in children in low-income countries including oral vaccine interference, negative impact of enteropathy and burden of enteric infections. However, when considering these alternative approaches the potential for impact on non-specific effects currently provided by an oral live attenuated rotavirus vaccine should be assessed [6,79].

Neonates have unique immune systems that are susceptible to a range of infections largely due to limited experience and exposure to microbes. The neonatal immunological milieu favours inhibition of T helper 1 (Th1) and pro-inflammatory responses that further impair innate and adaptive responses to neonatal diseases [65,101]. Furthermore, it has been shown that a high frequency of suppressive regulatory T cells that induce increased levels of the inhibitory cytokines IL-10 and TGF-β characterise the developing neonatal/infant immune system [102–104]. This inhibitory milieu reportedly limits the expansion of follicular Th1 cells and sequentially diminishes germinal centre B cell responses that are critical for generation of immune memory [65]. Studies of the immunomodulatory effects of live attenuated rotavirus vaccines, such as RV3-BB, on the neonatal immune system would allow us to gain understanding of the potential off-target effects of this novel and promising live attenuated vaccine. These effects might confer additional benefits over its protective effects against rotavirus infection with positive implications in vulnerable neonates and infants [105].
Figure 2: Illustration of potential beneficial off-target effects of live attenuated RV3-BB vaccine: At birth and in younger infants, the immune system generally favours inhibition of Th1 and innate responses (A). Live-attenuated vaccines such as RV3-BB administered at birth (B) would circumvent the imbalance towards enhanced immune responses (C) \cite{30,99,101}.

A lack of an appropriate research model to study off-target effects in neonates and infants has presented a challenge to further understanding. Human neonatal trials are often impractical and ethically challenging thereby limiting interrogation of immune responses in early life. Neonatal mouse models provide a practical means of investigating off-target effects and vaccine interventions, however, inherent variations between humans and mice restrains the direct extrapolation of findings to humans \cite{65}. Umbilical cord blood is an alternative means of assessing \textit{ex vivo} immune responses in human neonates prior to vaccination. Use of neonatal gnotobiotic pig (Gn) in biomedical research is yet another valuable model and is so far a proven model of human rotavirus disease \cite{81,106}. Pigs and humans share high genomic and protein sequence, thus facilitating extrapolation of findings to humans \cite{107}.

Umbilical cord blood samples can be leveraged to provide useful insights into the immune ontogeny and the suspected vaccine influence in inducing off-target effects of vaccines \cite{108}. A comparison between cord blood sample before and after vaccine intervention could uncover useful insights in the context of potential off-target effects induced by vaccines administered within the first days of life. It is therefore imperative that more mechanistic studies of off-target effects are conducted for live-attenuated rotavirus vaccines in order to generate evidence that could inform understanding of any beneficial off-target effects and lead to improved vaccines that are tailored to the unique neonatal and infant immune system.
Figure 3: Potential areas that require further study in the context of off-target effects of vaccination. Factors with potential to influence induction of off-target effects include geographical setting, sex/genetics, vaccine type, performance and vaccine co-administration, vaccine timing, composition of gut microbiome, prior maternal exposure to vaccine or microbes and disease burden in endemic settings. We hypothesise that a live attenuated rotavirus vaccine (RV3-BB) may cause the development of beneficial off-target vaccine effects when administered at birth, with the potential to interact with sex and infant gut microbiome contributing to overall reduced infant mortality.

Important mechanistic questions that require further attention include:

- How long do vaccine induced off-target effects last?
- To what extent do external factors such as maternal vaccine exposure, birth method, and vaccine type influence occurrence of off-target effects?
- What mechanistic approaches can separate vaccine induced off-target effects versus environmentally induced off-target effects (such as early pathogen exposure, infant gut microbiome, maternal vaccine/pathogen exposure)?
- At what age does sex influence the occurrence of off-target effects?
- Where adequate evidence on sex interaction with vaccine induced effects is gathered, what is the practicability of developing vaccines tailored to specific sex or about interindividual variations that occur naturally?
- In the context of a neonatal live attenuated RV3-BB vaccine, can co-administration of RV3-BB with known off-target inducers such as BCG affect induction of anticipated off-target effects?

As the hunt for improved knowledge and understanding of vaccine induced off-target effects continues, interpretation of findings must take into consideration potential cofounders such as the geographical or epidemiological setting, maternal vaccine or pathogen exposure, biological factors such as sex, disease burden to infections such as norovirus, cytomegalovirus and vaccine timing (Figure 3). Attributing changes in morbidity or mortality to vaccine effects is often challenging, especially in high mortality low income setting with concurrent and changing public health and social-welfare interventions. Therefore, robust study designs which use relevant quality data to accurately quantify indirect vaccine effects and overall vaccine impact is essential before estimating any off-target effects in the real world.

In summary, live-attenuated rotavirus vaccines have the potential to induce beneficial off-target effects which could be clinically significant when administered to the immunologically naïve neonate. The live-attenuated neonatal
rotavirus vaccine, RV3-BB, administered from birth offers potential to provide early protection against severe rotavirus disease but also infection from other pathogens via its off-target effects on the immune system of the newborn.

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**Conflicts of Interest:**

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**References**

[1] WHO. Children: improving survival and well-being n.d. https://www.who.int/news-room/fact-sheets/detail/children-reducing-mortality (accessed October 11, 2021).

[2] Clark A, Black R, Tate J, Roose A, Kotloff K, Lam D, et al. Estimating global, regional and national rotavirus deaths in children aged <5 years. PLoS One 2017;12:e0183392.

[3] WHO. Diarrhoeal disease n.d. https://www.who.int/news-room/fact-sheets/detail/diarrhoeal-disease (accessed October 11, 2021).

[4] WHO. Vaccines and immunization 2021. https://www.who.int/health-topics/vaccines-and-immunization#tab=tab_1 (accessed December 8, 2021).

[5] Cárcamo-Calvo R, Muñoz C, Buesa J, Rodríguez-Díaz J, Gozalbo-Rovira R. The rotavirus vaccine landscape, an update. Pathogens 2021;10:1–15. https://doi.org/10.3390/pathogens10050520.

[6] Burke RM, Tate JE, Kirkwood CD, Steele AD, Parashar UD. Current and new rotavirus vaccines. Curr Opin Infect Dis 2019;32:435–44. https://doi.org/10.1097/QCO.0000000000000572.

[7] Troeger et al Rotavirus Vaccination and the Global Burden of Rotavirus Diarrhea Among Children Younger
Than 5 Year n.d.

[8] Glass RI. Unexpected benefits of rotavirus vaccination in the United States. J Infect Dis 2011;204:975–7. https://doi.org/10.1093/infdis/jir477.

[9] Bar-Zeev N, King C, Phiri T, Beard J, Mvula H, Crampin AC, et al. Impact of monovalent rotavirus vaccine on diarrhoea-associated post-neonatal infant mortality in rural communities in Malawi: a population-based birth cohort study. Lancet Glob Heal 2018;6:e1036–44. https://doi.org/10.1016/S2214-109X(18)30314-0.

[10] Lopman BA, Curns AT, Yen C, Parashar UD. Infant rotavirus vaccination may provide indirect protection to older children and adults in the United States. J Infect Dis 2011;204:980–6. https://doi.org/10.1093/infdis/jir492.

[11] Uthayakumar D, Paris S, Chapat L, Freyburger L, Poulet H, De Luca K. Non-specific Effects of Vaccines Illustrated Through the BCG Example: From Observations to Demonstrations. Front Immunol 2018;9:2869. https://doi.org/10.3389/fimmu.2018.02869.

[12] Shann F. The nonspecific effects of vaccines and the expanded program on immunization. J Infect Dis 2011;204:182–4. https://doi.org/10.1093/infdis/jir244.

[13] Higgins JPT, Soares-weiser K, López-lópez JA, Kakourou A, Chaplin K, Christensen H, et al. Association of BCG, DTP , and measles containing vaccines with childhood mortality: systematic review n.d. https://doi.org/10.1136/bmj.2016.i5170.

[14] Kleinnijenhuis J, Quintin J, Preijers F, Joosten LAB, Ifrim DC, Saeed S, et al. Bacille Calmette-Guérin induces NOD2-dependent nonspecific protection from reinfection via epigenetic reprogramming of monocytes. Proc Natl Acad Sci U S A 2012;109:17537–42. https://doi.org/10.1073/pnas.1202870109.

[15] de Bree LCJ, Koeken VACM, Joosten LAB, Aaby P, Benn CS, van Crevel R, et al. Non-specific effects of vaccines: Current evidence and potential implications. Semin Immunol 2018;39:35–43. https://doi.org/10.1016/j.smim.2018.06.002.

[16] Shann F. Nonspecific Effects of Vaccines and the Reduction of Mortality in Children. Clin Ther 2013;35:109–14. https://doi.org/10.1016/j.clinthera.2013.01.007.

[17] WHO. Meeting of Global Advisory Committee on Vaccine Safety. Meet Glob Advis Comm Vaccine Saf n.d.;Weekly Epi:287–292.

[18] Pennington SH, Ferreira DM, Caamaño-Gutiérrez E, Reîné J, Hewitt C, Hyder-Wright AD, et al. Nonspecific effects of oral vaccination with live-attenuated Salmonella Typhi strain Ty21a. Sci Adv 2020;5:1–11. https://doi.org/10.1126/SCIADV.AAU6849.

[19] Aaby P, Benn CS, Flanagan KL, Klein SL, Kollmann TR, Lynn DJ, et al. The non-specific and sex-differential effects of vaccines. Nat Rev Immunol 2020;20:464–70. https://doi.org/10.1038/s41577-020-0338-x.

[20] Sankoh O, Welaga P, Debuur P, Zandoh C, Gyaase S, Poma MA, et al. The non-specific effects of vaccines and other childhood interventions: The contribution of INDEPTH Health and Demographic Surveillance Systems. Int J Epidemiol 2014;43:645–53. https://doi.org/10.1093/ije/dyu101.

[21] Aaby P, Martins CL, Garly ML, Balé C, Andersen A, Rodrigues A, et al. Non-specific effects of standard measles vaccine at 4.5 and 9 months of age on childhood mortality: Randomised controlled trial. BMJ 2010;341:c6495. https://doi.org/10.1136/bmj.c6495.

[22] Roth AE, Benn CS, Ravn H, Rodrigues A, Lisse IM, Yazdanbakhsh M, et al. Effect of revaccination with BCG in early childhood on mortality: Randomised trial in Guinea-Bissau. BMJ 2010;340:c671. https://doi.org/10.1136/bmj.c671.

[23] Moorlag SJCFM, Arts RJW, van Crevel R, Netea MG. Non-specific effects of BCG vaccine on viral infections. Clin Microbiol Infect 2019;25:1473–8. https://doi.org/10.1016/j.cmi.2019.04.020.

[24] De Groot AS, Ardito M, McClaine EM, Moise L, Martin WD. Immunoinformatic comparison of T-cell epitopes contained in novel swine-origin influenza A (H1N1) virus with epitopes in 2008-2009 conventional influenza
vaccine. Vaccine 2009;27:5740–7. https://doi.org/10.1016/j.vaccine.2009.07.040.

[25] Saadatian-Elahi M, Aaby P, Shann F, Netea MG, Levy O, Louis J, et al. Heterologous vaccine effects. Vaccine 2016;34:3923–30. https://doi.org/10.1016/j.vaccine.2016.06.020.

[26] Goodridge HS, Ahmed SS, Curtis N, Kollmann TR, Levy O, Netea MG, et al.Harnessing the beneficial heterologous effects of vaccination. Nat Rev Immunol 2016;16:392–400. https://doi.org/10.1038/nri.2016.43.

[27] Kleinnijenhuis J, Quintin J, Preijers F, Benn CS, Joosten LAB, Jacobs C, et al. Long-lasting effects of bcg vaccination on both heterologous th1/th17 responses and innate trained immunity. J Innate Immun 2014;6:152–8. https://doi.org/10.1159/000355628.

[28] Smith SG, Kleinnijenhuis J, Netea MG, Dockrell HM. Whole Blood Profiling of Bacillus Calmette–Guérin-Induced Trained Innate Immunity in Infants Identifies Epidermal Growth Factor, IL-6, Platelet-Derived Growth Factor-AB/BB, and Natural Killer Cell Activation. Front Immunol 2017;8:1–11. https://doi.org/10.3389/fimmu.2017.00644.

[29] Ritz N, Mui M, Balloch A, Curtis N. Non-specific effect of Bacille Calmette-Guérin vaccine on the immune response to routine immunisations. Vaccine 2013;31:3098–103. https://doi.org/10.1016/j.vaccine.2013.03.059.

[30] Kollmann TR, Kampmann B, Mazmanian SK, Marchant A, Levy O. Protecting the Newborn and Young Infant from Infectious Diseases: Lessons from Immune Ontogeny. Immunity 2017;46:350–63. https://doi.org/10.1016/j.immuni.2017.03.009.

[31] Brook B, Harbeson DJ, Shannon CP, Cai B, He D, Ben-Othman R, et al. BCG vaccination-induced emergency granulopoiesis provides rapid protection from neonatal sepsis. Sci Transl Med 2020;12. https://doi.org/10.1126/scitranslmed.AAX4517.

[32] Cuenca AG, Cuenca AL, Gentile LF, Efron PA, Islam S, Moldawer LL, et al. Delayed emergency myelopoiesis following polymicrobial sepsis in neonates. Innate Immun 2015;21:386–91. https://doi.org/10.1177/1753425914542445.

[33] Netea MG, Joosten LAB, Latz E, Mills KHG, Stunnenberg HG, Neill LAJO, et al. Trained immunity: A programme of innate immune memory in health and disease 2017;352:1–23. https://doi.org/10.1016/science.aaf1098 Trained.

[34] Schnack L, Sohrabi Y, Lagache SMM, Kahles F, Bruemmer D, Waltenberger J, et al. Mechanisms of trained innate immunity in oxLDL primed human coronary smooth muscle cells. Front Immunol 2019;10:1–12. https://doi.org/10.3389/fimmu.2019.00013.

[35] Naik S, Larsen SB, Gomez NC, Alaverdyan K, Yuan S, Polak L, et al. HHS Public Access 2018;550:475–80. https://doi.org/10.1038/nature24271.Inflammatory.

[36] Lu Y, Sun Y, Drummer C, Nanayakkara GK, Shao Y, Saaoud F, et al. Increased acetylation of H3K14 in the genomic regions that encode trained immunity enzymes in lysophosphatidylcholine-activated human aortic endothelial cells – Novel qualification markers for chronic disease risk factors and conditional DAMPs. Redox Biol 2019;24:101221. https://doi.org/10.1016/j.redox.2019.101221.

[37] Hamada A, Torre C, Drancourt M, Ghigo E. Trained immunity carried by non-immune cells. Front Microbiol 2019;10:1–8. https://doi.org/10.3389/fmicb.2018.03225.

[38] dos Santos JC, Barroso de Figueiredo AM, Teodoro Silva MV, Cirovic B, de Bree LCJ, Damen MSMA, et al. β-Glucan-Induced Trained Immunity Protects against Leishmania braziliensis Infection: a Crucial Role for IL-32. Cell Rep 2019;28:2659–2672.e6. https://doi.org/10.1016/j.celrep.2019.08.004.

[39] Milutinović B, Kurtz J. Immune memory in invertebrates. Semin Immunol 2016;28:328–42. https://doi.org/10.1016/j.smim.2016.05.004.

[40] Melillo D, Marino R, Italiani P, Boraschi D. Innate Immune Memory in Invertebrate Metazoans: A Critical Appraisal. Front Immunol 2018;9:1915. https://doi.org/10.3389/fimmu.2018.01915.
[41] Sohrabi Y, Dos Santos JC, Dorenkamp M, Findeisen H, Godfrey R, Netea MG, et al. Trained immunity as a novel approach against COVID-19 with a focus on Bacillus Calmette–Guérin vaccine: mechanisms, challenges and perspectives. Clin Transl Immunol 2020;9. https://doi.org/10.1002/cti2.1228.

[42] Bryant C, Kerr ID., Debnath M, Kenny KHA, Ratnam J, Ferreira RS., et al. Candida albicans Infection Affords Protection against Reinfected via Functional Reprogramming of Monocytes. Cell Host Microbe 2020;19:6218–21. https://doi.org/10.1016/j.chom.2012.06.006.Candida.

[43] Arts RJW, Moorlag SJCFM, Novakovic B, Li Y, Wang SY, Oosting M, et al. BCG Vaccination Protects against Experimental Viral Infection in Humans through the Induction of Cytokines Associated with Trained Immunity. Cell Host Microbe 2018;23:89-100.e5. https://doi.org/10.1016/j.chom.2017.12.010.

[44] Bai W, Liu H, Ji Q, Zhou Y, Liang L, Zheng R, et al. TLR3 regulates mycobacterial RNA-induced IL-10 production through the PI3K/AKT signaling pathway. Cell Signal 2014;26:942–50. https://doi.org/10.1016/j.cellsig.2014.01.015.

[45] Speth MT, Repnik U, Müller E, Spanier J, Kalinke U, Corthay A, et al. Poly(I:C)-Encapsulating Nanoparticles Enhance Innate Immune Responses to the Tuberculosis Vaccine Bacille Calmette-Guérin (BCG) via Synergistic Activation of Innate Immune Receptors. Mol Pharm 2017;14:4098–112. https://doi.org/10.1021/acs.molpharmaceut.7b00795.

[46] Aaby P, Roth A, Ravn H, Napirna BM, Rodrigues A, Lisse IM, et al. Randomized trial of BCG vaccination at birth to low-birth-weight children: Beneficial nonspecific effects in the neonatal period? J Infect Dis 2011;204:245–52. https://doi.org/10.1093/infdis/jir240.

[47] Biering-Sørensen S, Aaby P, Napirna BM, Roth A, Ravn H, Rodrigues A, et al. Small randomized trial among low-birth-weight children receiving bacillus Calmette-Guérin vaccination at first health center contact. Pediatr Infect Dis J 2012;31:306–8. https://doi.org/10.1097/INF.0b013e3182458289.

[48] Aaby P, Ravn H, Roth A, Rodrigues A, Lisse IM, Dines BR, et al. Early diphtheria-tetanus-pertussis vaccination associated with higher female mortality and no difference in male mortality in a cohort of low birthweight children: An observational study within a randomised trial. Arch Dis Child 2012;97:685–91. https://doi.org/10.1136/archdischild-2011-300646.

[49] Aaby P, Jensen H, Gomes J, Fernandes M, Lisse IM. The introduction of diphtheria-tetanus-pertussis vaccine and child mortality in rural Guinea-Bissau: An observational study. Int J Epidemiol 2004;33:374–80. https://doi.org/10.1093/ije/dyh005.

[50] Shann F. Commentary: BCG vaccination halves neonatal mortality. Pediatr Infect Dis J 2012;31:308–9. https://doi.org/10.1097/INF.0b013e3182465be8.

[51] Bucardo F, Reyes Y, Svensson L, Nordgren J. Predominance of norovirus and sapovirus in nicaragua after implementation of universal rotavirus vaccination. PLoS One 2014;9:1–8. https://doi.org/10.1371/journal.pone.0098201.

[52] Becker-Dreps S, Cuthbertson CC, Bucardo F, Vinje J, Paniagua M, Giebultowicz S, et al. Environmental factors associated with childhood norovirus diarrhoea in León, Nicaragua. Epidemiol Infect 2017;145:1597–605. https://doi.org/10.1017/S0950268817000322.

[53] Iturriza-Gómar M, Jere KC, Hungerford D, Bar-Zeev N, Shioda K, Kanjerwa O, et al. Etiology of Diarrhea among Hospitalized Children in Blantyre, Malawi, following Rotavirus Vaccine Introduction: A Case-Control Study. J Infect Dis 2019;220:213–8. https://doi.org/10.1093/infdis/jiz084.

[54] Burnett E, Parashar U, Tate J. Rotavirus Vaccines: Effectiveness, Safety, and Future Directions. Pediatr Drugs 2018;20:223–33. https://doi.org/10.1007/s40272-018-0283-3.

[55] Organizations WH, Record WE, States M, Group SA, Africa S. Rotavirus vaccines WHO position paper: January 2013 - Recommendations. Vaccine 2013;31:6170–1. https://doi.org/10.1016/j.vaccine.2013.05.037.
Aaby P, Benn C, Nielsen J, Lisse IM, Rodrigues A, Ravn H. Testing the hypothesis that diphtheria e tetanus e pertussis vaccine has negative non-specific and sex-differential effects on child survival in high-mortality countries 2012:1–24. https://doi.org/10.1136/bmjopen-2011-000707.

Aaby P, Benn CS. Developing the concept of beneficial non-specific effect of live vaccines with epidemiological studies 2019:25. https://doi.org/10.1016/j.cmi.2019.08.011.

Aaby P, Kollmann TR, Benn CS. C O M M E N T A R Y Nonspecific effects of neonatal and infant vaccination: public-health, immunological and conceptual challenges 2014:895–900.

Aaby P, Jensen H, Samb B, Cisse B, Sodemann M, Jakobsen M, et al. Differences in female-male mortality after high-titre measles vaccine and association with subsequent vaccination with diphtheria-tetanus-pertussis and inactivated poliovirus: reanalysis of West African studies 2003;361:2183–9.

Aaby P, Garly M.G, Nielsen J RH. Increased Female-Male Mortality Ratio Associated With. Pediatr Infect Dis J 2007;26:247–52. https://doi.org/10.1097/01.inf.0000256735.05098.01.

Benn CS, Fisker AB, Rieckmann A, Sorup S, Aaby P. Personal View Vaccinology: time to change the paradigm? n.d.:274–83. https://doi.org/10.1016/S1473-3099(19)30742-X.

Klein SL, Shann F, Moss WJ, Benn CS, Aabye P. RTS,S malaria vaccine and increased mortality in girls. Mbio 2016;7:1–2. https://doi.org/10.1128/mBio.00514-16.

WHO. Meeting of the strategic Advisory Group of Experts on Immunization. Wkly Epidemiol Rec 2014;21:221–36.

Noho-Konteh F, Adetifa JU, Cox M, Hossin S, Reynolds J, Le MT, et al. Sex-Differential Non-Vaccine-Specific Immunological Effects of Diphtheria-Tetanus-Pertussis and Measles Vaccination. Clin Infect Dis 2016;63:1213–26. https://doi.org/10.1093/cid/ciw492.

Saso A, Kampmann B. Vaccine responses in newborns. Semin Immunopathol 2017;39:627–42. https://doi.org/10.1007/s00281-017-0654-9.

Flanagan KL, Klein SL, Skakkebaek NE, Marriott I, Marchant A, Selin L, et al. Sex differences in the vaccine-specific and non-targeted effects of vaccines. Vaccine 2011;29:2349–54. https://doi.org/10.1016/j.vaccine.2011.01.071.

Klein SL, Flanagan KL. Sex differences in immune responses. Nat Rev Immunol 2016;16:626–38. https://doi.org/10.1038/nri.2016.90.

Flanagan KL, Fink AL, Plebanski M, Klein SL. Sex and gender differences in the outcomes of vaccination over the life course. Annu Rev Cell Dev Biol 2017;33:577–99. https://doi.org/10.1146/annurev-cellbio-100616-060718.

Fink AL, Engle K, Ursin RL, Tang WY, Klein SL. Biological sex affects vaccine efficacy and protection against influenza in mice. Proc Natl Acad Sci U S A 2018;115:12477–82. https://doi.org/10.1073/pnas.1805268115.

Potluri T, Fink AL, Sylvia KE, Dhakal S, Vermillion MS, vom Steeg L, et al. Age-associated changes in the impact of sex steroids on influenza vaccine responses in males and females. Npj Vaccines 2019;4. https://doi.org/10.1038/s41541-019-0124-6.

Bishop R.F, Barnes G.L, Ciprian E LJ. Clinical Immunity after Neonatal Rotavirus Infection — A Prospective Longitudinal Study in Young Children. N Engl J Med 1983;309:72–6.

Simonsen L, Viboud C, Elixhauser A, Taylor RJ, Kapikian AZ. More on RotaShield and intussusception: The role of age at the time of vaccination. J Infect Dis 2005;192. https://doi.org/10.1086/431512.

Rennels M, Parashar U, Holman R, Le C, Chang H GR. Lack Of An Apparent Association Between Intussusception and wild or vaccine Rotavirus infection. Pediatr Infect Dis J 1998;17:924–5.

Arnold MM. Rotavirus Vaccines: Why Continued Investment in Research Is Necessary. Curr Clin Microbiol Reports 2018;5:73–81. https://doi.org/10.1007/s40588-018-0079-9.

Grant L, Watt J, Moulton L, Weatherholtz R, Reid R, Santosham M, et al. Lack of nonspecific protection against
all-cause nonrotavirus gastroenteritis by vaccination with orally administered rotavirus vaccine. J Pediatr Gastroenterol Nutr 2013;56:635–40. https://doi.org/10.1097/MPG.0b013e318287c5ec.

[76] King C, Bar-Zeev N, Phiri T, Beard J, Mvula H, Crampin A, et al. Population impact and effectiveness of sequential 13-valent pneumococcal conjugate and monovalent rotavirus vaccine introduction on infant mortality: prospective birth cohort studies from Malawi. BMJ Glob Heal 2020;5:e002669. https://doi.org/10.1136/bmjgh-2020-002669.

[77] Bergman H, Henschke N, Hungerford D, Pitan F, Ndwandwe D, Cunliffe N, et al. Vaccines for preventing rotavirus diarrhoea: vaccines in use. Cochrane Database Syst Rev 2021;2021. https://doi.org/10.1002/14651858.CD008521.pub6.

[78] Clarke E, Desselberger U. Correlates of protection against human rotavirus disease and the factors influencing protection in low-income settings. Mucosal Immunol 2015;8:1–17. https://doi.org/10.1038/mi.2014.114.

[79] Bines JE, Kotloff KL. Next-generation rotavirus vaccines: important progress but work still to be done. Lancet Infect Dis 2020;20:762–4. https://doi.org/10.1016/S1473-3099(20)30151-1.

[80] Fix A, Kirkwood CD, Steele D, Flores J. Next-generation rotavirus vaccine developers meeting: Summary of a meeting sponsored by PATH and the bill & melinda gates foundation (19–20 June 2019, Geneva). Vaccine 2020;38:8247–54. https://doi.org/10.1016/j.vaccine.2020.11.034.

[81] Bishop RF, Tzipori SR, Coulson BS, Unicomb LE, Albert MJ, Barnes GL. Heterologous protection against rotavirus-induced disease in gnotobiotic piglets. J Clin Microbiol 1986;24:1023–8. https://doi.org/10.1128/jcm.24.6.1023-1028.1986.

[82] Bines JE, Danchin M, Jackson P, Handley A, Watts E, Lee KJ, et al. Safety and immunogenicity of RV3-BB human neonatal rotavirus vaccine administered at birth or in infancy: A randomised, double-blind, placebo-controlled trial. Lancet Infect Dis 2015;15:1389–97. https://doi.org/10.1016/S1473-3099(15)00227-3.

[83] Segovia MJG. Human neonatal rotavirus vaccine (RV3-BB) to target rotavirus from birth. Acta Pediatr Esp 2018;76:62–3. https://doi.org/10.1056/nejmoa1706804.

[84] Boniface K, Byars SG, Cowley D, Kirkwood CD, Bines JE. Human neonatal rotavirus vaccine (RV3-BB) produces vaccine take irrespective of histo-blood group antigen status. J Infect Dis 2020;221:1070–8. https://doi.org/10.1093/infdis/jiz333.

[85] Pollock L, Bennett A, Jere KC, Dube Q, Mandolo J, Bar-Zeev N, et al. Nonsecretor Histo-blood Group Antigen Phenotype Is Associated with Reduced Risk of Clinical Rotavirus Vaccine Failure in Malawian Infants. Clin Infect Dis 2019;69:1313–9. https://doi.org/10.1093/cid/ciy1067.

[86] Sharma S, Hagbom M, Svensson L, Nordgren J. The impact of human genetic polymorphisms on rotavirus susceptibility, epidemiology, and vaccine take. Viruses 2020;12. https://doi.org/10.3390/v12030324.

[87] Witte DA. Neonatal Rotavirus Vaccine (RV3-BB) immunogenicity and safety in a neonatal and infant administration schedule in Malawi: A randomized, double-blind, four-arm parallel group dose-ranging study. Prepr with Lancent 2021;Available.

[88] Witte D, Handley A, Jere KC, Bogandovic-Sakran N, Mpakiza A, Turner A, et al. Neonatal Rotavirus Vaccine (RV3-BB) Immunogenicity and Safety in a Neonatal and Infant Administration Schedule in Malawi:&nbsp;A Randomized, Double-Blind, Four-Arm Parallel Group Dose-Ranging Study. SSRN Electron J 2021;3099:1–11. https://doi.org/10.2139/ssrn.3858564.

[89] Bar-Zeev N, Kapanda L, Tate JE, Jere KC, Iturriza-Gomara M, Nakagomi O, et al. Effectiveness of a monovalent rotavirus vaccine in infants in Malawi after programmatic roll-out: An observational and case-control study. Lancet Infect Dis 2015;15:422–8. https://doi.org/10.1016/S1473-3099(14)71060-6.

[90] Mandolo JJ, Henrion MYR, Mhango C. Reduction in severity of all-cause gastroenteritis requiring hospitalisation in children vaccinated against rotavirus in Malawi 2021;1–23.
[91] Hungerford DJ, French N, Iturriza-Gómez M, Read JM, Cunliffe NA, Vivancos R. Reduction in hospitalisations for acute gastroenteritis-associated childhood seizures since introduction of rotavirus vaccination: A time-series and change-point analysis of hospital admissions in England. J Epidemiol Community Health 2019;73:1020–5. https://doi.org/10.1136/jech-2019-213055.

[92] Lee YJ, Lee JY, Jang YH, Seo SU, Chang J, Seong BL. Non-specific effect of vaccines: Immediate protection against respiratory syncytial virus infection by a live attenuated influenza vaccine. Front Microbiol 2018;9:1–11. https://doi.org/10.3389/fmicb.2018.00083.

[93] Seo SU, Lee KH, Byun YH, Kweon MN, Seong BL. Immediate and broad-spectrum protection against heterologous and heterotypic lethal challenge in mice by live influenza vaccine. Vaccine 2007;25:8067–76. https://doi.org/10.1016/j.vaccine.2007.09.012.

[94] Mohanty S, Joshi SR, Ueda I, Wilson J, Blevins TP, Siconolfi B, et al. Prolonged proinflammatory cytokine production in monocytes modulated by interleukin 10 after influenza vaccination in older adults. J Infect Dis 2015;211:1174–84. https://doi.org/10.1093/infdis/jiu573.

[95] Lanthier PA, Huston GE, Moquin A, Eaton SM, Szaba FM, Kummer LW, et al. Live attenuated influenza vaccine (LAIV) impacts innate and adaptive immune responses. Vaccine 2011;29:7849–56. https://doi.org/10.1016/j.vaccine.2011.07.093.

[96] Koyama S, Ishii KJ, Kumar H, Tanimoto T, Coban C, Uematsu S, et al. Differential Role of TLR- and RLR-Signaling in the Immune Responses to Influenza A Virus Infection and Vaccination. J Immunol 2007;179:4711–20. https://doi.org/10.4049/jimmunol.179.7.4711.

[97] Lund JM, Alexopoulou L, Sato A, Karow M, Adams NC, Gale NW, et al. Recognition of single-stranded RNA viruses by Toll-like receptor 7. Proc Natl Acad Sci U S A 2004;101:5598–603. https://doi.org/10.1073/pnas.0400937101.

[98] Xue Q, Tang Y, Huang G. Innate immune responses in RNA viral infection. Front Med 2020. https://doi.org/10.1007/s11684-020-0776-7.

[99] Kelvin AA, Zambon M. Influenza imprinting in childhood and the influence on vaccine response later in life. Eurosurveillance 2019;24:1–5. https://doi.org/10.2807/1560-7917.ES.2019.24.48.1900720.

[100] Freyne B, Messina NL, Donath S, Germano S, Bonnici R, Gardiner K, et al. Neonatal BCG Vaccination Reduces Interferon-γResponsiveness to Heterologous Pathogens in Infants from a Randomized Controlled Trial. J Infect Dis 2020;221:1999–2009. https://doi.org/10.1093/infdis/jiaa030.

[101] Dowling DJ, Levy O. Ontogeny of early life immunity. Trends Immunol 2014;35:299–310. https://doi.org/10.1016/j.it.2014.04.007.

[102] Morris MC, Surendran N. Neonatal vaccination: Challenges and intervention strategies. Neonatology 2016;109:161–9. https://doi.org/10.1159/000442460.

[103] Burt TD. Fetal Regulatory T Cells and Peripheral Immune Tolerance In Utero: Implications for Development and Disease. Am J Reprod Immunol 2013;69:346–58. https://doi.org/10.1111/aji.12083.

[104] Boer MC, Joosten SA, Ottenhoff THM. Regulatory T-cells at the interface between human host and pathogens in infectious diseases and vaccination. Front Immunol 2015;6:1–15. https://doi.org/10.3389/fimmu.2015.00217.

[105] Pulendran B, S. Arunachalam P, O'Hagan DT. Emerging concepts in the science of vaccine adjuvants. Nat Rev Drug Discov 2021;20:454–75. https://doi.org/10.1038/s41573-021-00163-y.

[106] Twitchell EL, Tin C, Wen K, Zhang H, Becker-Dreps S, Azcarate-Peril MA, et al. Modeling human enteric dysbiosis and rotavirus immunity in gnotobiotic pigs. Gut Pathog 2016;8:1–18. https://doi.org/10.1186/s13099-016-0136-y.

[107] Wang M, Donovan SM. Human microbiota-associated swine: Current progress and future opportunities. ILAR J 2015;56:63–73. https://doi.org/10.1093/ilar/ilv006.
[108] Tokuhara D, Hikita N. Cord blood-based approach to assess candidate vaccine adjuvants designed for neonates and infants. Vaccines 2021;9:1–14. https://doi.org/10.3390/vaccines9020095.