Appendix 2 - Planned analyses

1 Baseline

Descriptive statistics:

For eligible children visited by a study nurse within 72 hours after birth, we will describe reasons for exclusion by group allocation.

Distribution of background factors will be presented by group allocation overall and by sex and region. Background factors will be summarised by counts (percentages), means (standard deviation) or medians (interquartile range) as appropriate. Information on the proportion with missing information will be provided.

Table 1: Summary of background factors by intervention and control group

- Sex
- Age at enrolment
- Region
- Weight at enrolment
- Temperature
- Mid-upper-arm circumference
- Head circumference
- Place of birth
- Socioeconomic factors (maternal education and housing conditions)

2 Primary analysis of primary outcome

The primary analysis of early infant non-accidental mortality will be assessed on a per-protocol (PP) analysis allowing for different baseline hazards according to factors used in the randomization (Region, pre-study mortality level (high/low)) and sex, thus allowing different baseline hazards for boys and girls. To account for clustering we will employ cluster-robust variance estimates. For the primary outcome, we will use Cox proportional hazards models that allow different baseline hazards according to the above mentioned factors and with age as underlying time-scale. Deaths due to accidents will be censored.

The primary analysis of the primary outcome is described in more detail in table 2.

Table 2: Primary analysis of primary outcome

Population	Children visited within 72 hours after birth are eligible for the study.
	Exclusion criteria:

Observation period	 Children already BCG vaccinated Moribund children (Expected not to survive the next 24 hours, as evaluated by the health facility nurse at the enrolment visit) Children in rural villages where the BHP mobile teams coincidentally were in the village the same day (and vaccinated all children) From: Enrolment visit or 24 hours after birth, whichever comes last
	To: 60 days of life
	Censoring, first of:
	- Visit by the BHP
	- 60 days
	- Death due to accident
	- Date of registering first non-trial vaccine given after
	enrolment
	- Migration (migration out of the study area as per HDSS
	definition)
Time scale	Age
Failure definition	Death
Statistical tool	Cox proportional hazards model
Stratification	We will employ models that allow for different baseline hazards
	according to sex, region and pre-study mortality level
Clustering	We will use robust standard errors to account for clustering within
	village cluster
Outline stata code	
For analysis:	stset outdate, f(dead==1) enter (max(dob+1, date_enrol) ///
	exit (min (dnasc+60, date_mobileteam) origin(dob)
	stcox random, strata(sex reg prmorlev) vce(cl regam)
For model checking:	estat phtest, detail†
	stphplot, strata(random) adj(sex reg prmorlev)
	stcox random, strata(sex reg prmorlev) vce(cl regam) tvc(random)
	texp(_t)
	*reg=region, prmorlev=pre-study mortality level, regam=village cluster
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[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

3 Effect-modifier analyses of primary outcome

We will assess whether the effect of the intervention on the primary effect measure is modified by the following potential effect modifiers.

Potential effect	Sex
modifier	
Design	We will perform the analysis as describe above (for the primary
	analysis) allowing the effect of the intervention to differ between the
	sexes.
Reasoning	Previous studies have found sex-differential non-specific effects ¹² ,
	therefore, we will assess the sex-differential effects.
Outline stata code:	
For analysis:	stcox random#sex sex, strata(sex reg prmorlev) vce(cl regam)
	contrast random#sex
For model checking:	estat phtest, detail†
	stphplot, strata(random_sex) adj(sex reg prmorlev)
	stcox random#sex sex, strata(sex reg prmorlev) vce(cl regam) ///
	<pre>tvc(random#sex sex) texp(_t)</pre>
	*reg=region, prmorlev=pre-study mortality level, regam=village
	cluster, random_sex=a four-level variable based on the four possible
	combinations of random and sex

Table 3. Sex as a potential effect modifier of the primary outcome

[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

Potential effect	Maternal BCG scar (yes/no)
modifier	
Design	We will perform the analysis as described above (for the primary
	analysis) allowing the effect of the intervention to differ by maternal
	BCG-scar status
Reasoning	A recent randomized trial in Denmark found that the effect of BCG
	varied by whether the mother had received BCG or not ³ . Since, BCG
	scar is a life-long marker of a successful BCG-vaccination, we will
	assess whether the effect of BCG differs by maternal BCG-scar status
Outline stata code:	
For analysis:	stcox random#mBCGscar mBCGscar, strata(sex reg prmorlev) vce(cl
	regam)

Table 4. Maternal BCG scar as a potential effect modifier of the primary outcome

	contrast random#mBCGscar
For model checking:	
	estat phtest, detail†
	stphplot, strata(random_mBCGscar) adj(sex reg prmorlev)
	stcox random#mBCGscar mBCGscar, strata(sex reg prmorlev) ///
	vce(cl regam) tvc(random#mBCGscar mBCGscar) texp(_t)
	* mBCGscar= maternal BCG scar, reg=region, prmorlev=pre-study mortality level, regam=village cluster, random_mBCGscar=a four- level variable based on the four possible combinations of random and maternal BCG scar

[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

Potential effect	Low birthweight (<2500g: yes/no)
modifier	
Design	We will perform the analysis as described above (for the primary
	analysis) allowing the effect of the intervention to differ by birthweight
	strata
Reasoning	Birthweight is an important risk factor for mortality. Previous
	randomised trials from Guinea-Bissau assessing the effect of BCG on
	mortality have been performed among low-birth-weight children.
Outline stata code:	
For analysis:	stcox random#LBW LBW, strata(sex reg prmorlev) vce(cl regam)
	contrast random#LBW
For model checking:	estat phtest, detail†
	stphplot, strata(random_LBW) adj(sex reg prmorlev)
	stcox random#LBW LBW, strata(sex reg prmorlev) vce(cl regam) ///
	tvc(random#LBW LBW) texp(_t)
	* LBW=Low birthweight, reg=region, prmorlev=pre-study mortality
	level, regam=village cluster, random_LBW=a four-level variable
	based on the four possible combinations of random and low
	birthweight

Table 5. Low birthweight as a potential effect modifier of the primary outcome

[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

Table 6. Season as a potential effect modifier of the primary outcome

Potential effect

Season of birth (Dry: December-May/Rainy: June-November)

i otentiai enteet	Season of birth (Dry. December-May/Ramy. June-November)
modifier	
Design	We will perform the analysis as described above (for the primary
	analysis) allowing the effect of the intervention to differ by season of
	birth
Reasoning	Previous studies have found that the effect of some vaccines is stronger
	in the dry season ⁴ . Therefore, we would like to assess if the effect of
	BCG and OPV differs according to season.
Outline stata code:	
For analysis:	stcox random#season season, strata(sex reg prmorlev) vce(cl regam)
	contrast random#season
For model checking:	estat phtest, detail [†]
	stphplot, strata(random_season) adj(sex reg prmorlev)
	stcox random#season season, strata(sex reg prmorlev) vce(cl regam) ///
	tvc(random#season season) texp(_t)
	*reg=region, prmorlev=pre-study mortality level, regam=village
	cluster, random_season=a four-level variable based on the four
	possible combinations of random and season

[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

4 Primary analyses of secondary outcomes

Non-accident morbidity

Since it can be difficult to distinguish between hospital admissions and outpatient contact through interviews, we have defined hospital admissions as an overnight stay in a health facility. Hospital admissions due to accidents are ignored but the follow-up time is (interval) censored while the child is admitted.

Table 7: Non-accident hospitalisation

Population	Identical to primary analysis of primary outcome
Observation period	From: Enrolment visit or 24 hours after birth, whichever comes last
	To: 60 days of life
	Censoring, first of:
	- Visit by the BHP
	- Date of registering first non-trial vaccine given after
	enrolment
	- Death
	- Hospital admission due to accident

	- 60 days
	 Migration (migration out of the study area as per HDSS definition)
Time scale	Age
Failure definition	First hospital admission – only overnight hospitalisations or arrival at
	the hospital and death within the first day will be considered in this
	analysis.
Statistical tool	Cox proportional hazards model
Stratification	We will employ models that allow for different baseline hazards
	according to sex, region and pre-study mortality level
Clustering	We will use robust standard errors to account for clustering within
	village cluster
Outline stata code	
For analysis:	stset outdate, f(hosp==1) enter (max(dob+1, date_enrol) ///
	exit (min (dnasc+60, date_mobileteam) origin(dob)
	stcox random, strata(sex reg prmorlev) vce(cl regam)
For model checking:	estat phtest, detail†
	stphplot, strata(random) adj(sex reg prmorlev)
	stcox random, strata(sex reg prmorlev) vce(cl regam) tvc(random)
	texp(_t)
	*reg=region, prmorlev=pre-study mortality level, regam=village cluster

[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

Severe morbidity

We will evaluate the effect on severe morbidity considered as the composite outcome of nonaccidental death and non-accidental hospital admission. Since it can be difficult to distinguish between hospital admissions and outpatient contact through interviews, we have defined hospital admissions as an overnight stay in a health facility. Hospital admissions due to accidents are ignored but the follow-up time is (interval) censored while the child is admitted. The potential effect modifiers for the primary outcome specified in tables 3-6 will also be assessed for the composite outcome.

Table 8: Severe morbidity

Population	Identical to primary analysis of primary outcome
Observation period	From: Enrolment visit or 24 hours after birth, whichever comes last

	To: 60 days of life
	Censoring, first of:
	- Visit by the BHP
	- Date of registering first non-trial vaccine given after
	enrolment
	- Death/Hospital admission due to accident
	- 60 days
	- Migration (migration out of the study area as per HDSS definition)
Time scale	Age
Failure definition	Death or first hospital admission
Statistical tool	Cox proportional hazards model
Stratification	We will employ models that allow for different baseline hazards
	according to sex, region and pre-study mortality level
Clustering	We will use robust standard errors to account for clustering within
	village cluster
Outline stata code:	
For analysis	stset outdate, f(event==1) enter (max(dob+1, date_enrol) ///
	exit (min (dnasc+60, date_mobileteam) origin(dob)
	stcox random, strata(sex reg prmorlev) vce(cl regam)
For model checking	estat phtest, detail†
	stphplot, strata(random) adj(sex reg prmorlev)
	stcox random, strata(sex reg prmorlev) vce(cl regam) tvc(random)
	texp(_t)
	*reg=region, prmorlev=pre-study mortality level, regam=village cluster
+ If the manage and a set have a	rds assumption is not fulfilled we will still report the marginal hazard ratio

[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

Table 9: All-cause consultations

Population	Identical to primary analysis of primary outcome
Observation period	From: Enrolment visit or 24 hours after birth, whichever comes last To: 60 days of life
	Censoring, first of: - Visit by the BHP
	 Date of registering first non-trial vaccine given after enrolment 60 days

	 Migration (migration out of the study area as per HDSS definition) Death End of study 	
Failure definition	An out-patient consultation within the observation period	
Statistical tool	Log-binomial regression	
Adjustment	We will adjust the analysis for sex, region and pre-study mortality level	
Clustering	We will use robust standard errors to account for clustering within village cluster	
Stata code	Binreg cons random sex b1.reg prmorlev, rr vce(cl regam) *cons=out-patient consultation, reg=Region, prmorlev=Pre-study mortality level, regam=village cluster	

Growth

Table 10: Mid-upper-arm circumference (MUAC)

Population	Identical to primary analysis of primary outcome	
Observation time point	First visit by the mobile teams	
Growth measures	MUAC will be analysed using the measured value	
Statistical tool	Linear regression	
Adjustment	We will adjust the analysis for MUAC at enrolment, age at MUAC	
	measurement, sex, region and pre-study mortality level	
Clustering	We will use robust standard errors to account for clustering within	
	village cluster	
Stata code	Regress MUAC random sex b1.reg prmorlev MUACenrol	
	MUACage, /// vce(cl regam)	
	*reg=region, prmorlev=pre-study mortality level,	
	MUACenrol=MUAC at enrolment, MUACage= age at MUAC	
	assessment, regam=village cluster	

Table 11: Weight-for-age z-score

Population	Identical to primary analysis of primary outcome	
Observation time point	First visit by the mobile teams	
Growth measures	Weight will be analysed using the WHO weight-for-age z-score	
Statistical tool	Linear regression	
Adjustment	We will adjust the analysis for weight-for-age at enrolment, sex,	
	region and pre-study mortality level	
Clustering	We will use robust standard errors to account for clustering within	
	village cluster	

Stata code	Regress w-z-score random w-z-enrol sex b1.reg prmorlev, vce(cl
	regam)
	*w-z-score=weight-for-age z-score at first visit by the mobile teams,
	w-z-enrol=weight-for-age z-score at enrolment, reg=region,
	prmorlev=pre-study mortality level, regam=village cluster

BCG scarring

Table	12:	BCG	scarring
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Population	Identical to primary analysis of primary outcome	
Observation timepoint	First visit after 6 months of age	
Failure definition	Scar (yes/no)	
Statistical tool	Log-binominal regression	
Adjustment	We will adjust the analysis for sex, region and pre-study mortality	
	level	
Clustering	We will use robust standard errors to account for clustering within	
	village cluster	
Stata code	Binreg scar random sex b1.reg prmorlec, rr vce(cl regam)	
	*reg=region, prmorlev=pre-study mortality level, regam=village	
	cluster	

Cost-effectiveness of providing BCG and OPV at birth

A cost effectiveness analysis seeking to measure the cost per death averted using a societal perspective will be performed, contrasting the costs of vaccine provision in the present programme and an outreach system as tested in the trial. The costs/savings associated with different rates of consultations and admissions will also be taken into account.

Suppurative lymphadenitis

We will assess the incidence of suppurative lymphadenitis as a reaction to BCG vaccination in the intervention and control clusters. Other serious adverse events to the BCG and OPV vaccine will be captured through the outcome measures (mortality, hospital admission and consultations).

5 Sensitivity analyses to test for robustness of conclusions

Table 13: Cause-specific death

Population	Identical to primary analysis of primary outcome	
Observation period	From: Enrolment visit or 24 hours after birth, whichever comes last	
	To: 60 days of life	
	Censoring, first of:	

	- Visit by the BHP	
	- 60 days	
	- Death due to accident	
	- Date of registering first non-trial vaccine given after	
	enrolment	
	- Migration (migration out of the study area as per HDSS	
	definition)	
Time scale	Age	
Failure definition	Death due to:	
	Malaria, Respiratory Infection, Sepsis, Gastrointestinal disease, Other	
Statistical tool	Cox proportional hazards model	
Stratification	We will employ models that allow for different baseline hazards	
	according to sex, region and pre-study mortality level	
Clustering	We will use robust standard errors to account for clustering within	
	village cluster	
Outline stata code		
For analysis:	stset outdate, f(event==1&cause==X) enter (max(dob+1, date_enrol)	
	/// exit (min (dnasc+60, date_mobileteam) origin(dob)	
	stcox random, strata(sex reg prmorlev) vce(cl regam)	
For model checking:	estat phtest, detail†	
	stphplot, strata(random) adj(sex reg prmorlev)	
	stcox random, strata(sex reg prmorlev) vce(cl regam) tvc(random)	
	texp(_t)	
	*reg=region, prmorlev=pre-study mortality level, regam=village	
	cluster	
+ If the propertional hazar	ds assumption is not fulfilled we will still report the marginal hazard ratio	

[†] If the proportional hazards assumption is not fulfilled we will still report the marginal hazard ratio but it will be supplemented by hazard ratios for two-three categorized time periods.

In sensitivity analyses, we will furthermore, assess whether the conclusions are robust to the following:

- Censoring follow-up at general health intervention campaigns (e.g. OPV campaigns)
- Altering the population to using an intention-to-treat approach, including all children, who had a home visit by a study nurse (i.e., including children who did not receive assigned treatment, children who were not enrolled because they were moribund, or who did not accept to participate).

- Altering the outcome from non-accidental mortality to all-cause mortality including also deaths due to accidents.
- For the non-accident hospitalisations, we will perform the analyses allowing for repeated hospitalisations. A child will return to the at-risk population the day after discharge from the hospital.

References

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- 3. Stensballe LG, Ravn H, Birk NM, et al. BCG Vaccination at Birth and Rate of Hospitalization for Infection Until 15 Months of Age in Danish Children: A Randomized Clinical Multicenter Trial. *J Pediatric Infect Dis Soc* 2018;0(0):1-8. doi: 10.1093/jpids/piy029 [published Online First: 2018/04/11]
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