Longitudinal antibody and T cell responses in Ebola virus disease survivors and contacts: an observational cohort study

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Summary

Background The 2013–16 Ebola virus disease epidemic in west Africa caused international alarm due to its rapid and extensive spread resulting in a significant death toll and social unrest within the affected region. The large number of cases provided an opportunity to study the long-term kinetics of Zaire ebolavirus-specific immune response of survivors in addition to known contacts of those infected with the virus.

Methods In this observational cohort study, we worked with leaders of Ebola virus disease survivor associations in two regions of Guinea, Guéckédou and Coyah, to recruit survivors of Ebola virus disease, contacts from households of individuals known to have had Ebola virus disease, and individuals who were not knowingly associated with infected individuals or had not had Ebola virus disease symptoms to serve as negative controls. We did Zaire ebolavirus glycoprotein-specific T cell analysis on peripheral blood mononuclear cells (PBMCs) on location in Guinea and transported plasma and PBMCs back to Europe for antibody quantification by ELISA, functional neutralising antibody analysis using live Zaire ebolavirus, and T cell phenotype studies. We report on the longitudinal cellular and humoral response among Ebola virus disease survivors and highlight potentially pauciymptomatic infection.

Findings We recruited 117 survivors of Ebola virus disease, 66 contacts, and 23 negative controls. The mean neutralising antibody titre among the Ebola virus disease survivors 3–14 months after infection was 1/174 (95% CI 1/136–1/223). Individual results varied greatly from 1/10 to more than 1/1000 but were on average ten times greater than that induced after 1 month by single dose Ebola virus vaccines. Following reactivation with glycoprotein peptide, the mean T cell responses among 116 Ebola virus disease survivors as measured by ELISpot was 305 spot-forming units (95% CI 257–353). The dominant CD8+ polyfunctional T cell phenotype, as measured among 53 Ebola virus disease survivors, was interferon γ+, tumour necrosis factor+, interleukin-2–, and the mean response was 0·046% of total CD8+ T cells (95% CI 0·021–0·071). Additionally, both neutralising antibody and T cell responses were detected in six (9%) of 66 Ebola virus disease contacts. We also noted that four (3%) of 137 individuals with Ebola virus disease infections did not have circulating Ebola virus-specific antibodies 3 months after infection.

Interpretation The continuous high titre of neutralising antibodies and increased T cell response might support the concept of long-term protective immunity in survivors. The existence of antibody and T cell responses in contacts of individuals with Ebola virus disease adds further evidence to the existence of sub-clinical Ebola virus infection.

Funding US Food & Drug Administration, Horizon 2020 EU EVIDENT, Wellcome, UK Department for International Development.

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Introduction

Dogma suggests that Ebola virus disease survivors have long-term protection against reinfection with Zaire ebolavirus and studies in non-human primates and rodents indicate that such protective effects are mediated to a great extent by both antibody and T cell responses to the viral envelope surface glycoprotein.1–4 Vaccine studies in non-human primates also suggest that CD8+, interferon (IFN)+, tumour necrosis factor+, interleukin 2 (IL-2)+/+ T cells are a potential correlate of protection.1–5,6 The Ebola virus vaccine, Ervebo, which is based on the viral glycoprotein, was evaluated during the 2013–16 epidemic. Ring vaccination trials estimated vaccine efficacy to be 100%.7 Previous studies on survivors of Ebola virus disease immunology have been done but only antibody levels were monitored.8 During historical Ebola virus outbreaks, plasma from survivors had been used to treat individuals suffering from acute Ebola virus
Research in context

Evidence before this study

We searched the PubMed database with MESH terms “Ebola,” “immunity,” “sero-prevalence,” “paucisymptomatic,” and for articles published between database inception and April 15, 2020, in any language. There was one report on long-term antibody and T cell responses in 11 survivors split across two small Sudan ebolavirus outbreaks. Neutralisation assays revealed titres of 0–1/80, and T cell proliferation and kinetics analyses clearly showed CD8+ T cell activation, but quantification by ELISpot was not done. A comprehensive analysis of ELSA responses in survivors from the 1995 Kikwit outbreak showed antibody titres plateaued at 3 weeks to 1 year after onset of symptoms. A report by Leroy and colleagues revealed the existence of paucisymptomatic individuals with Ebola virus infections. Additionally, work by Glynn and colleagues further demonstrates the presence of asymptomatic Ebola virus infection. Several reports showed persistence of Ebola virus in immune privileged sites, and an in-depth longitudinal analysis of the B cell response among four Ebola virus disease survivors has also been done. We found no other studies reporting the longitudinal analysis of cellular and humoral immunity among survivors over 3 consecutive years.

Added value of this study

This unique study reports an in-depth analysis of naturally acquired immunity to Ebola virus and enables a comprehensive comparison between naturally acquired and vaccine-induced immunity to Ebola virus both at the antibody and T cell level. The study also provides supporting evidence for the existence of paucisymptomatic Ebola virus disease and suggests that true incidence of Ebola virus infection in the west African outbreak was greater than recorded. Additionally, T cell phenotyping results support the preclinical findings of a potential correlate of protection to Ebola virus.

Implications of all the available evidence

We suggest that up to 9% of individuals with Ebola virus disease will present with mild symptoms, which will have implications with regard to monitoring and responding to future outbreaks. Additionally, the work here shows that T cell responses are relevant and long lasting among survivors of Ebola virus disease; therefore, future vaccine developers might wish to consider the T-cell response in more depth. The evaluation of convalescent plasma showed varying titres of anti-Ebola virus IgG, so pre-screening of IgG and neutralising titres before administration of convalescent plasma should be considered in future outbreaks.

Methods

Study designs and participants

We did an observational study based on opportunistic sampling. Volunteers were recruited from two prefectures in Guinea, Guéckédou, the epicentre of the outbreak, and Coyah, which is around 300 miles away. Blood was collected from survivors of Ebola virus disease 3–14 months after infection. A second and third bleed of volunteers was done about 12 and 24 months later. Individuals presented their Ebola virus disease survivor certificate or were identified on Ebola treatment centre databases to verify that they were survivors. Contacts were defined as individuals who provided care for or were living in the same household of an Ebola virus disease confirmed case. Survivors and contacts were asked a number of questions (appendix 2, pp 2–4) regarding their contact with individuals with Ebola virus disease, as well as presentation of any Ebola virus disease-like symptoms.
Following immune analysis, where possible, a more in-depth interview took place to understand the degree of care they gave and discuss symptoms they might have had. Additionally, volunteers were recruited, who were not knowingly exposed to people with Ebola virus disease and did not attend high-risk events such as funerals. All volunteers were informed of the procedures and purpose of the study and only consenting participants were included. Ethical approval was obtained from the National Ethics Committee for Health Research, Guinea (33/CNERS/15) and from the National Research Ethics Service, UK. Written, informed consent was obtained from all participants.

**Procedures**

30–50 mL of blood was collected into edetic acid vacutainers. Plasma and peripheral blood mononuclear cells (PBMCs) were separated by layering the blood over a Ficoll gradient (GE Healthcare; Amersham, UK) and centrifugation. Plasma was aspirated and stored at –20°C for antibody studies. PBMCs were either used fresh, or frozen (–70°C) and shipped to the UK, where they were stored in liquid nitrogen. Participants were interviewed using the health surveillance questionnaire (appendix 2, pp 2–4) and with regards to health scores during the Ebola outbreak each category was given equal weighting with a maximum score of 13. Freshly isolated PBMCs were prepared at 2×10⁶ cells/mL in Leibovitz media supplemented with penicillin–streptomycin, foetal calf serum, L-glutamine, HEPES, and 2-mercaptoethanol was used for IFNy ELISPot. Briefly, whole blood was layered onto LeucoSep-tubes (Sigma; Gillingham, UK) containing Ficoll-Paque Plus (GE Healthcare; Amersham, UK); the sample was centrifuged at 800 G for 20 min and PBMCs were collected from the buffy coat layer. The PBMCs were stimulated with Ebola virus glycoprotein peptide library (Mimotopes; Melbourne, Australia) at a final concentration of 2.5 µg per peptide per mL, as described previously.⁷ After 18–20 h incubation at 37°C, IFNy release was determined by standard ELISPot protocol (Mabtech; Nacka Strand, Sweden) and spot forming cells enumerated using an S6 core analyser (Cellular Technology; Shaker Heights, OH). IFNy release was calculated by subtracting the background from each well and taking the mean of three triplicate wells. The results were determined as spot forming units per one million cells and IFNy response to the Ebola virus glycoprotein peptide were summed to determine the overall T-cell response.

ELISA was done as described previously.⁸ Briefly, high binding microtitre plates were coated with whole Ebola virus inactivated virions and incubated for 16–20 h. After washing in PBS and 0·1% Tween20 (PBST) and blocking (PBS and 5% milk powder), 1/200 dilutions of plasma sample were added to the plates and incubated for 1 h. Horseradish peroxidase-conjugated polyclonal antibody (P0214, Dako; Santa Clara, CA; dilution 1/1000) in conjunction with TMB substrate was used to develop the reaction. Optical density was determined at 450 nm minus 630 nm (reference wavelength). Each sample was analysed in duplicate on mock and viral antigen. The mean optical density of each sample on the mock antigen was subtracted from the mean optical density of the respective sample on the Ebola virus antigen. Arbitrary ELISA units were extrapolated by linear regression analysis using standard curves generated from patient antiserum. Further specificity was assessed using a Zaire, Makona Ebola virus glycoprotein (sourced from Nuffield Department of Medicine, Oxford University, Oxford, UK) specific ELISA, for which Nunc Maxisorb 96-well plates (Merck; Darmstadt, Germany) were coated overnight (16–18 h) with purified Ebola virus glycoprotein antigen (0·5 µg/mL). Plasma was serially diluted, starting at 1/200 and the bound IgG was detected using goat anti-human IgG Fcγ specific antibody conjugated to alkaline phosphatase (1/15000). Alkaline phosphatase-yellow substrate (Sigma) was added and the optical density measured at 405 nm using a VERSAmax plate reader (Molecular Devices; Wokingham, UK) controlled by SoftMax Pro Enterprise software (version 4.71). The plates were read using a predefined Softmax template, which fits a four-parameter logistic curve to the dose response data. The cutoff was defined as the mean negative value plus five SDs.

The activity of the Ebola virus-specific antibodies present in plasma was determined by neutralisation of Ebola virus variant Mayinga (1976) as previously described.⁹ Briefly, after heat treatment for complement inactivation, plasma was serially diluted in supplemented Dulbecco’s modified Eagle’s medium (DMEM) in 96-well culture plates, 100 median tissue culture infectious dose (TCID₅₀) units of Ebola virus variant Mayinga were added to the plasma dilutions. Following incubation at 37°C for 1 h, Vero cell suspension in supplemented DMEM was added. Plates were then incubated at 37°C with 5% CO₂ and cytopathic effects were evaluated at 7 days after infection. Neutralisation titres were calculated as geometric mean titre of four replicates. A titre of 1/8 or above is classified as positive.

Recombinant glycoprotein, nucleoprotein, and viral protein 35 were generated based on the Ebola virus strain Makona in HEK293T cells and whole cell lysates were used. Viral protein 40 was based on Ebola virus strain Kikwit and was obtained from Stratech Scientific (Ely, UK). Proteins were heat denatured and loaded onto 4–12% BisTris gels and separated by size by SDS-PAGE. The proteins were then transferred to polyvinylidene difluoride membrane and blocked overnight in block buffer (PBST buffer with 5% milk). Plasma was diluted 1/1000 in block buffer and incubated with the Ebola virus-protein containing blots for 4 h at room temperature. The blots were washed for 5 min in PBST. Secondary antibody (goat anti-human IgG [γ-chain...
specific peroxidase conjugate) F(ab’)2 fragments (Sigma; A2290), were diluted 1/1000 in block buffer. The blots were incubated with secondary antibody for 1 h at room temperature. Membranes were washed and blots developed with ECL prime, incubating for 5 min. Images were captured at 5 min and 10 min exposure and presence of immunoreactivity determined against a molecular marker standard.

Intracellular cytokine staining was done as has been described previously. Briefly, PBMCs were resuspended in warmed complete media (Roswell Park Memorial Institute medium supplemented with penicillin–streptomycin, fetal calf serum, L-glutamine, HEPES, and 2-mercaptoethanol) and rested overnight at 37°C. The following day cells were adjusted to 1 x 10⁶ cells/mL in media containing anti-CD28 BVU737, CD49d, and CD107a-PerCP cy5.5 (1 µg/mL). Samples were then left either untreated or were stimulated with an Ebola virus glycoprotein peptide pool, containing 187 15mer overlapping peptides at 2.5 µg/peptide or 1 µg/mL staphylococcal enterotoxin B for 16–18 h, as previously described. 2 h into the incubation, brefeldin A and monensin (1 µg/mL) were added to block cytokine secretion from the cell. The following day samples were washed in cold FACS wash and LIVE/DEAD fixable aqua dye (Life Technologies; Carlsbad, CA) was added. Samples were washed, then incubated with a cell surface cocktail of antibodies including CD3-APC 750, CD4-BV786, CD8-AF700, CD19-BV510, CD14-BV510, CCR7-APC, CD95-BUV395, and CD45RO-BV605. Cells were then washed, fixed and permeabilised using Becton Dickinson (London, UK) Cytofix/Cytoperm, and stained for intracellular cytokines using IFNγ-AF488, TNF-BV421, and IL-2-PE. Samples were then washed, resuspended, and acquired using a Becton Dickinson Fortessa machine and FACS Diva software. Sample analysis used FlowJo, Pestle, and SPICE software as described previously.1 All antibodies were obtained from Biolegend (London, UK), with the exception of CD95-BUV395 and CD28 BVU737, which were obtained from Becton Dickinson.

Statistical analysis
The data collected from all the volunteers were categorised into three groups, survivor, contact, and negative, which were sub-divided by region and sex. Measurements for ELISPot, ELISA and neutralisation were tested independently. Statistical analysis on the fixed effect coefficient for year was done using R version 4.0.1 with the lme4 package version 1.1–23 and fitted models were assessed for violation of assumptions. Cytokine responses were determined by subtracting the untreated response from that of the stimulation; negative values were set to 0-001 and statistical differences were determined using Mann-Whitney test. Correlations were determined using Spearman correlation analysis. Statistical tests were done using GraphPad Prism version 8.

Role of the funding source
The funder of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results
206 volunteers were recruited from the Guéckédou and Coyah prefectures (table; appendix 2, p 1): 117 survivors of Ebola virus disease, 66 contacts, and 23 negative controls. Glycoprotein-specific IgG titres were significantly correlated with whole Ebola virus-specific antibody titres (r 0.85; p < 0.0001; appendix 2, p 6). 113 (96%) of the 117 survivors had detectable concentrations of IgG to Ebola virus 3–14 months after infection, with responses measured at one dilution and defined as arbitrary units. 11 (17%) of 66 contacts had concentrations of Ebola virus-specific IgG that were above background (figure 1A). The activity of the Ebola virus-specific IgG was measured by the ability to neutralise the infection of Ebola virus-Mayinga in vitro. Testing for functional activity using our neutralisation assay showed that 113 (96%) of 117 survivors showed detectable titres of Ebola virus-specific neutralising antibody (figure 1B). Notably, there was a greater than 100-fold range in neutralisation titres (1/10–1/10000) among survivors, with a mean of 1/174. Plasma from six (9%) of the 66 contacts also showed Ebola virus-specific neutralising antibodies, and the mean of these six samples was 1/133, which paralleled their detectable concentrations of Ebola virus glycoprotein-specific IgG. Further characterisation of five of these samples by Western blot revealed the presence of antibodies not only to glycoprotein but also to nucleoprotein, viral protein 40, and viral protein 35 (appendix 2, p 7). All negative controls had undetectable neutralising activity. PBMCs from 15 (13%) of 116 survivors and isolated 4 months after infection did not produce an IFNγ response following glycoprotein peptide stimulation. During 2015 sampling, ELISpots were only done on contacts in the Coyah cohort, with the exception of G012 and G034 from the Guéckédou cohort; however, of the 42 contacts tested, eight had a detectable IFNγ response to Ebola virus glycoprotein. Six of these also had detectable Ebola virus-specific ELISA and neutralisation titres. Ebola virus disease survivors typically had a score of 7–5 out of 13 in health surveillance questionnaires, whereas close contacts would report a score of 1–3 out of 13. Of the contacts that showed a humoral and cellular response, only two of the five we were able to question reported a score greater than 2 out of 13, suggesting they were seropositive contacts that were displaying more overt symptoms during the outbreak.

We were able to longitudinally map the responses of 96 survivors and found that ELISA (figure 2A), neutralisation (figure 2B), and ELISPOT responses (figure 2C) appeared stable over time. Mixed-effects linear regression for each outcome showed that there was
statistical evidence for the effect of year for all three measured outcomes (assessed by ANOVA against nested intercept-only models; ELISA $p<0.0001$, neutralisation $p<0.0001$, ELISpot $p=0.0019$). Comparison of each model to a reduced model with a single linear parameter fitted for year showed that the linear predictor sufficiently explained the annual variance in the outcome for ELISA data only ($p=0.14$). For the ELISA outcome, the fixed effect coefficient for year showed an inverse relationship ($−1022.08$ arbitrary ELISA units, 95% CI $−1453.54$ to $−590.62$), suggesting an annual decrease in antibody titre. However, for all outcomes, the magnitude of serological responses across years and individuals remained higher than responses measured among control group samples (figure 1). These prolonged, elevated, responses could theoretically be due to re-exposure to Ebola virus antigen or simply reflect the basal immunological state to such a pathogenic infection. To address this issue, we plotted 2015 survivor results in relation to their time since resolution of infection. Results were consistent over time (figure 2D).

Flow cytometry and intracellular cytokine staining showed that the majority of IFNγ, as a proportion of their

|                | Guéckédou | Coyah | Total |
|----------------|-----------|-------|-------|
|                | Survivor  | Contact| Negative| Survivor| Contact| Negative| Survivor| Contact| Negative|
| Total          | 46        | 26    | 9      | 71      | 40    | 14      | 117      | 66     | 23      |
| Male           | 21        | 7     | 8      | 32      | 23    | 9       | 53       | 30     | 17      |
| Female         | 25        | 19    | 1      | 39      | 17    | 5       | 64       | 36     | 6       |
| Returning in 2016–17 | 33    | 19    | 0      | 63      | 23    | 0       | 96       | 42     | 0       |

Table: Numbers of participants from Coyah and Guéckédou sample sites

Figure 1: Characterisation of immune responses of Ebola virus disease survivors and close contacts
(A) Ebola virus-specific antibody titre against whole virus preparation as measured by ELISA in plasma samples collected from survivors (n=117), contacts (n=66), and negative controls (n=23). (B) Geometric mean neutralising antibody titre against live Ebola virus (strain Mayinga) using plasma samples collected from survivors (n=117), contacts (n=66), and negative controls (n=23). Results displayed on a log2 scale. (C) ELISpot to detect IFNγ secretion using an Ebola virus glycoprotein peptide library and peripheral blood mononuclear cell samples from survivors (n=116), contacts (n=42), and negative controls (n=21). (D) Quantified results from health questionnaire given to all survivors and contacts. A score of 13 represents severe symptoms and 0 represents no symptoms during the 2013-16 outbreak. Bars represent the geometric mean plus or minus the standard error of the mean. Dashed lines represent the mean plus five SDs. Mann-Whitney tests were done for all statistical analyses. Blue dots represent seronegative survivors. Coloured squares represent six (9%) of the 66 contacts who had Ebola virus-specific neutralising antibodies. IFNγ=interferon γ.
parent subset, was coming from CD8+ T cells (figure 3). CD8+ T cells originating from Ebola virus disease survivors (n=51) produced significantly more IFNγ than peripheral blood mononuclear cells originating from ELISpot negative controls (n=26; p<0·0001). Furthermore, when these IFNγ responses were backgated to their parent phenotype markers, the majority of antigen-specific CD8+ T cells showed intermediate CCR7 expression and low CD45RO expression, corresponding with a naive-like phenotype. By contrast, CD4+ T cells showed primarily CD45RO positive and CCR7 low or intermediate expression consistent with a central memory or effector memory phenotype.

A previous report suggested a role for so-called polyfunctional CD8+ T cells, which produce multiple cytokines, in the control of Ebola virus disease; we therefore investigated the incidence of IFNγ+, TNF+ or IFNγ+, TNF+, IL-2+ CD8+ T cells among 53 survivor PBMC samples collected in 2016 (15–28 months after infection). When these PBMCs were stimulated overnight...
with Ebola virus glycoprotein peptide pools, a significant proportion of T cells were IFNγ+, TNF+, and CD8+ compared with 26 contacts who were seronegative by ELISA and neutralisation (p=0·0004; figure 4). Significant populations of IFNγ+ (p=0·018) only and IFNγ+, TNF+ (p=0·0075) CD8+ T cells were identified among the survivors’ PBMCs compared with ELISpot negative controls. Survivor samples had a significantly larger proportion of CD107a expressing cells than their comparative contacts (p=0·032), and when backgated, the majority of these CD107a+ cells were IFNγ+, TNF+, CD8+ T cells (figure 4B). CD8+ T cells originating from survivors of Ebola virus disease showed significantly greater amounts of CD107a than negative controls (p=0·032). There was a strong correlation between total Ebola virus ELISA and neutralisation assay results (appendix 2, p 8). We also observed a significant correlation between ELISpot and ELISA results (appendix 2, p 8). We did not find a significant correlation or difference between either age, sex, or virus load at time of diagnosis (cycle threshold values) and any of the immunological parameters measured (appendix 2, pp 8–9).

**Discussion**

We report the most comprehensive study of Ebola virus disease survivor immunity to date and uniquely assess both neutralising antibody and T cell responses. Neutralising antibody titres from survivors 3–35 months after infection were ten-fold larger than those seen 1 month after vaccination with an efficacious single dose vaccine5,7. This finding is compelling evidence for long-term protection against reinfection with Ebola virus and bodes well for survivors if the disease returns to west Africa, suggesting that they could continue to have a role in front-line activities to control future outbreaks. However, the absence of humoral immunity in a small percentage of survivors might explain observations that a minority of Ebola virus disease survivors can be reinfected, suggesting it would be essential to assess their immunity before the risk of potential re-exposure. Of the four survivors who displayed no antibody response, two showed a detectable ELISpot response; however, this response was not present in subsequent samples and misuse of survivor certificates cannot be ruled out in this instance. Absence of antibody responses has been documented for various emerging diseases, including severe acute respiratory syndrome coronavirus 2 antibody negative samples from COVID-19 convalescent individuals.19 From these data we would suggest that sero-epidemiology studies for Ebola virus might underestimate the prevalence of the disease. The disparity between vaccinees and survivors also highlights the need to periodically boost vaccine immunity or use alternative heterologous vaccine vector approaches that might induce a more durable response. However, it is yet to be
Articles

The continuously high titres of neutralising antibody and T cell responses are perhaps unsurprising, because Ebola virus has been reported to persist for more than 1 year in immune privileged sites. Furthermore, where we have found responses to be stable up to 3 years after infection, studies of survivors from Ebola virus disease outbreaks suggest that ELISA antibody responses continue to be relatively stable for more than 1 year and that low titres of neutralising antibodies have been found in survivors 40 years after infection. However, further work is needed to show antigen persistence in these survivor samples.

We found steady state titres of Ebola virus glycoprotein antibody measured by ELISA and neutralisation from groups of survivors 3–14 months after the onset of symptoms. This finding contrasts with a study of four Ebola virus disease survivors in which longitudinal analysis suggest a continuous increase in neutralising titre from up to 900 days after recovery in some individuals. Obvious study differences, including an absence of longitudinal sampling and of experimental treatment, might explain the differing results. Of note, the longitudinal analysis in our study uses only data from participants who provided a full 3 years’ worth of data; therefore, data from participants with missing or incomplete data was discarded, which is a limitation of this study with regard to the stability over time. Long-term, potent Ebola virus responses could be a result of continuous exposure to viral antigen as it seeps back into the systemic environment from immune privileged locations. There is evidence that relapse and transmission can occur long after the primary infection has been resolved, and it would be of great interest to sample such immune privileged sites from our cohorts to see if there is live virus present. However, such an approach was not possible owing to biosafety constraints and the limitations of our ethical approval. We suggest that our studies do not support the presence of recirculating antigen 15 months or more after recovery, because we believe that CD8+ T cell phenotyping revealed a possible subset of T memory stem cells. T memory stem cells, are a self-renewing population of lymphocytes that are CD45RO-, CCR7+, CD27+, and CD95+. In support of this suggestion, the phenotype of IFNγ-producing CD4+ T cells was CD45RO+ (figure 3D), suggesting that these cells are memory T cells and not terminally differentiated effectors that have recently been exposed to antigen. LaVergne et al found that post Ebola syndrome was associated with greater activation of CD4+ and CD8+ T cells, so it would be of interest to correlate the magnitude of our T cell responses with post Ebola syndrome in any follow up studies.

Plasma from survivors of Ebola virus disease, containing various levels of neutralising antibodies, has been used to treat acute cases of Ebola virus infection during several prior outbreaks. However, the 2013–16 outbreak enabled a more comprehensive study to be done, which reported an absence of statistically significant improvement in survival associated with plasma treatment. In the same trial, neutralising antibody titres of plasma harvested from Ebola virus disease survivors were on average ten times smaller than seen in our study. However, differences between assay conditions could account for this outcome, which might have affected the subsequent analysis of functional antibody activity and clinical outcome. Furthermore, Sahr et al showed a positive effect with plasma therapy in a small clinical trial in Sierra Leone.

Immune analysis of contacts of individuals with Ebola virus disease, who did not have a confirmed positive PCR test or report with typical disease to a treatment centre, revealed that up to 9% have both detectable humoral and cellular immunity. This finding could be explained by the existence of cross-reactive antibodies to Ebola virus proteins, a hypothesis that has been suggested by a previous study that analysed serum from German citizens and found a number of filovirus specific seropositive cases. However, our seropositive contacts showed both neutralising antibody and T cell responses, and we were able to show that five individuals also had antibodies specific to Ebola virus nucleoprotein, viral protein 35, and viral protein 40, thus confirming that they must have
come into contact with the virus. The extent to which these contacts were symptomatic remains unknown and is difficult to address after the event and largely relies on anecdotal evidence. Matters are further complicated owing to the stigma surrounding Ebola virus disease in west Africa, meaning that individuals who were infected at the time were likely to have downplayed their symptoms or not sought treatment. However, owing to the health questioning and interviews that took place we are confident that the paucisymptomatic contacts within our cohort did have mild disease that did not prevent them from conducting their normal duties. Therefore, these observations strongly suggest that four potential outcomes from Ebola virus exposure exist: symptomatic infection resulting in death, symptomatic infection with survival and immunity, paucisymptomatic infection with survival and immunity, and no infection and no immunity. Additional support for the existence and potential transmission from paucisymptomatic people comes from a report whereby a mother was able to transmit Ebola virus to her child through breastmilk. Both parents reported no symptoms of Ebola virus disease and blood samples were PCR negative; however, semen and breast milk samples were shown to be PCR positive, and breastfeeding was concluded to be the most likely route of transmission. Furthermore, a number of serological studies have suggested the existence of paucisymptomatic infection in west Africa. It might not be possible to categorically show that these positive contacts were paucisymptomatic or showed mild disease; however, this study and others suggest that this aspect of Ebola virus transmission and the role it might have in spreading Ebola virus disease needs to be considered during any future outbreaks. Additionally, the number of individuals infected with Ebola virus could be significantly larger than the officially reported numbers, which will also affect the extent of the reported mortality rate.

We show here that survivors of Ebola virus disease have a long-lasting T cell response. The T cell responses in our survivor cohort, measured some years after recovery from disease, are similar to that seen 6 months after vaccination with chimpanzee adenovirus and modified vaccinia Ankara boost. The importance of this T cell activity to Ebola virus vaccine design has been the subject of debate. The ChAd3, MVA boost regime has been shown to induce a strong T cell response and primates have revealed that CD8+ polyfunctional T cells might have a key role in protection from Ebola virus challenge, following vaccination with an adenovirus vector. Our results show that the IFNγ+ and TNFα+ double-positive CD8+ T cell population, which is known to be involved in long-term protection in primates, is also the dominant CD8+ T cell population in responses to restimulation with glycoprotein peptide in Ebola virus disease survivors. Therefore, further work should be done to determine the activity and proportion of CD8+ polyfunctional T cells in response to Ebola virus vaccination.

Contributors
PM, M3C, SG, and MKK conceived the study and experimental design with input from GB, MM, LL, MG, DP, OS, IK, EN, SB, HR, JH, AMH-R, and NM. Sample collection within Guinea was led by JAB and MKK with assistance from M3C, BMK, SK, and ARA. Sample processing in Guinea was led by RT, TT, YH, JAB, and PM with help from FRK, KS, AV, VV, BA, LO, AB, SD, and EN. Analysis of samples in the UK was done by RT, TT, YH, and KSR. Processing of whole virus ELISA and live virus neutralisation was led by TS with help from SKF and VK. The manuscript was primarily written by RT, TT, YH, VK, M3C, SI, JM, CM-F, SB, and SG with input from all authors.

Declaration of interests
SG received grants from European Commission and German Research Foundation for the conduct of this study. All other authors declare no competing interests.

Data sharing
De-identified individual participant data that underlie the results reported in this article and the study protocol will be made available to researchers who provide a methodologically sound proposal beginning 3 months and ending 5 years following publication. Proposals should be directed to miles.carroll@phe.gov.uk, and to gain access, data requestors will need to sign a data access agreement.

Acknowledgments
This work was funded by the US Food & Drug Administration (HHSP232201510104C) and Horizon 2020 EU’s EVIDENT research initiative (666100). TS received funds from the German Research Foundation (197785619/SFB 1021). CM-Z and SG were supported by the German Research Foundation (GU 883/3–1 and MU 3565/3–1). MWC is joint funded by Wellcome and the UK Department for International Development (214626/Z/18/Z). We would like to acknowledge the long-term support and commitment of Coyah and Guéckédou Ebola virus disease survivors associations and all the participants in this study. We are also extremely grateful to the Guinean authorities and members of Center for Training and Research on Priority Diseases including Malaria in Guinea, including Saidou Koyate, Abdoulaye Barry, and Ousmane Soumah for their support of this study. We would also like to acknowledge the tremendous cooperation of the personnel of the Coyah and Guéckédou Prefectural Health Directorates and Offices of the regional Prefects.

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